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(54) Title: OUTER MEMBRANE PROTEIN OF EHRLICHIA CANIS AND EHRLICHIA CHAFFEENSIS

(57) Abstract

The present invention relates to diagnostic tools for veterinary and human use which are used for serodiagnosing ehrlichiosis in mammals, particularly in members of the Canidae family and in humans. The present invention also provides polynucleotides which encode the outer membrane proteins of E. chafeensis. The polynucleotides encode an OMP-1 family of proteins of E. chafeensis and P30 family of proteins of E. canis. The present invention also provides the following isolated proteins of E. chafeensis OMP-1, OMP-1A, OMP-1B, OMP-1C, OMP-1D, OMP-1B, OMP-1B, OMP-1T, OMP-

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OUTER MEMBRANE PROTEIN OF EHRLICHIA CANIS AND EHRLICHIA CHAFFEENIS

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BACKGROUND OF THE INVENTION

The ehrlichiae are obligate intracellular bacteria that infect circulating leucocytes. Ehrlichia chafeensis infects the monocytes and macrophages in humans and causes human monocytic ehrlichiosis. The clinical manifestations of ehrlichiosis in humans are nonspecific and similar to Rocky Mountain spotted fever. The clinical manifestations include fever, chills, headache mylagia or vomiting and weight loss. Most patients have a history of tick exposure.

Ehrlichia canis infects and causes ehrlichiosis in animals belonging to the family Canidae. Canine chrlichiosis consists of an acute and a chronic phase. The acute phase is characterized by fever, serous nasal and ocular discharges, anorexia, depression, and loss of weight. The chronic phase is characterized by severe pancytopenia, epistaxis, hematuria, blood in feces in addition to more severe clinical signs of the acute disease. If treated early during the course of the disease, dogs respond well to doxycycline. However, chronically infected dogs do not respond well to the antibiotic. Therefore, early diagnosis is very important for treating canine ehrlichiosis.

The primary diagnostic test for diagnosing canine ehrlichiosis and human ehrlichiosis is the indirect fluorescent antibody (IFA) test. This test uses the etiologic agent Ehrlichia canis to diagnose canine ehrlichiosis. The IFA test uses Ehrlichia chafeensis as antigen for diagnosing human ehrlichiosis. The IFA test has, however, serious limitations. The IFA test is subject to false positives because the antigens are made of whole infected cells which comprise many nonspecific proteins which will cross-react with sera from some patients. The IFA test is also subject to false negatives because IFA antigens are unstable and may become inactivated during storage. In addition the IFA test requires a special equipment to perform the test. For example, the IFA test requires a tissue culture system for growing the bacterium that are used to prepare the antigen slides, a fluorescent microscope, and trained persons to evaluate the serum reactivity to the bacterial antigen on the slide.

Tools which permit simpler, more rapid, and objective serodiagnosis of canine chrlichiosis or human chrlichiosis are desirable.

SUMMARY OF THE INVENTION

The present invention relates to improved diagnostic tools for veterinary and human use which are used for serodiagnosing ehrlichiosis in mammals, particularly in members of the Canidae family and in humans.

The present invention also provides polynucleotides or nucleic acids which encode the outer membrane proteins of E. chafeensis. The OMP-1 polynucleotide encodes an OMP-1 protein of E. chafeensis having a molecular weight of about 27.7 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG.3B, SEQ ID NO: __. The OMP-1B polynucleotide encodes an OMP-1B protein of E.

chafeensis having a molecular weight of about 28.2 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 4B, SEQ ID NO: __. The OMP-1C polynucleotide encodes an OMP-1C protein of E. chafeensis having a molecular weight of about 27.6 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 5B, SEQ ID NO: __. The OMP-1D polynucleotide encodes an OMP-1D protein of E. chafeensis having a molecular weight of about 28.7 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 6B, SEQ ID NO: __. The OMP-1E polynucleotide encodes an OMP-1E protein of E. chafeensis having a molecular weight of about 27.8 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 7B, SEQ ID NO: __. The OMP-1F polynucleotide encodes an OMP-1F protein of E. chafeensis having a molecular weight of about 27.9 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 8B, SEQ ID NO: __. The OMP-1A polynucleotide encodes an OMP-1A protein of E. chafeensis having a molecular weight of about 29.6 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 9B, SEQ ID NO: ___. The OMP-1R polynucleotide encodes an OMP-1R protein of E. chafeensis having a molecular weight of at least 23 kDa and comprising an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 10B, SEQ ID NO: __. The OMP-1S polynucleotide encodes an OMP-1S protein of E. chafeensis comprising an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 11B, SEQ ID NO: ___ The OMP-1T polynucleotide encodes an OMP-1T protein of E. chafeensis comprising an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 12B, SEQ ID NO: __. The OMP-1U polynucleotide encodes an OMP-1U protein of E. chafeensis having a molecular weight of about 30.6 kDa and an amino acid sequence which is at least 85% homologous to amino acid sequence shown in FIG. 13B, SEQ ID NO: __. The OMP-1V polynucleotide encodes an OMP-1V protein of E. chafeensis having a molecular weight of about 28.0 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 14B, SEQ ID NO: __. The OMP-1W polynucleotide encodes an OMP-1W protein of E. chafeensis having a molecular weight of about 28.8 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 15B, SEQ ID NO: __. The OMP-1X polynucleotide encodes an OMP-1S protein of E. chafeensis having a molecular weight of about 27.8 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 16B, SEQ ID NO: __. The OMP-1Y polynucleotide encodes an OMP-1Y protein of E. chafeensis having a molecular weight about 28.8 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 17B, SEQ ID NO: __. The OMP-1Z polynucleotide encodes an OMP-1Z protein of E. chafeensis comprising an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 18B, SEQ ID NO: ___.

The outer membrane proteins from E. chaffeensis, particularly a recombinant form of OMP-1, are immunogenic and, thus are useful for preparing antibodies. Such antibodies are useful for immunolabeling isolates of E. chafeensis and for detecting the presence of E. chafeensis in body fluids, tissues, and particularly in monocytes and macrophages. The isolated outer membrane proteins, particularly OMP-1, are also useful for

membrane protein, particularly OMP-1, are also useful immunogens for raising antibodies that are capable of reducing the level of infection in an immunized mammal that has been infected with E. chafeensis. The isolated membrane proteins are also useful in a vaccine for protecting against infection with E. chafeensis.

The present invention also relates to isolated polynucleotides which encode 30 kDa outer membrane proteins from Ehrlichia canis. The proteins are designated P30 and P30a. The proteins, particularly P30, are immunogenic and are, thus, useful for preparing antibodies that are useful for immunolabeling isolates of E. canis. The P30 protein is also useful for diagnosing canine ehrlichiosis in mammals, particularly in members of the family Canidae, most particularly in dogs and for diagnosing infections with E. chafeensis in humans. The P30 protein is also a useful immunogen for raising antibodies that reduce the level of infection in an immunized mammal that has been infected with E. canis. The P30 protein is also useful in a vaccine for protecting animals against infection with E. canis.

The present invention also provides the following isolated proteins of E. chafeensis OMP-1 (also known as p28), OMP-1A, OMP-1B, OMP-1C, OMP-1D, OMP-1E, OMP-1F, OMP-1R, OMP-1S, OMP-1T, OMP-1U, OMP-1V, OMP-1W, OMP-1X, and OMP-1Z, referred to hereinafter collectively as the "OMP family". The present invention also provides the following isolated proteins of E. canis P30, P30-a, P30-1, P30-2, P30-3, P30-4, P30-5, P30-6, P30-7, P30-8, P30-9, and P30-10, referred to hereinafter as the P30 family.

The present invention also relates to an assay for diagnosing ehrlichiosis in humans using a recombinant outer membrane protein of E. chafeensis, particularly OMP-1. The present invention also relates to an assay for diagnosing ehrlichiosis in humans and members of the family Canidae using a recombinant outer membrane protein of E. canis, particularly P30.

Brief Description of the Figures

- FIG. 1. shows the DNA sequence of and the amino acid sequence encoded by the *E. chafeensis* (p28) gene cloned in pCRIIp28. The N-terminal amino acid sequence of native omp-1 protein (P28) determined chemically is underlined. Five amino acid residues at the N terminus of P28 which were not included in the p28 gene, are indicated by boldface. Arrows indicate annealing positions of the primer pair designed for PCR
- FIG. 2. shows the restriction map of 6.3-kb genomic DNA including the *omp-1* gene copies in *E. chafeensis*. The four DNA fragments were cloned from the genomic DNA (pPS2.6, pPS3.6, pEC2.6, and pEC3.6). A recombinant plasmid pPS2.6 has an overlapping sequence with that of pEC3.6. The closed boxes at the bottom show PCR-amplified fragments from the genomic DNA for confirmation of the overlapping area. Open boxes at the top indicate open reading frames (ORF) of *omp-1* gene copies with direction by arrows. Open boxes at the bottom show DNA fragments subcloned for DNA sequencing.
- FIG. 3B shows one embodiment of the OMP-1 protein; FIG. 3A shows one embodiment of the OMP-1 polynucleotide.
- FIG. 4B shows one embodiment of the OMP-1B protein, FIG. 4A shows one embodiment of the OMP-1B polynucleotide

FIG. 5A shows one embodiment of the OMP-1C polynucleotide; FIG 5B shows one embodiment of the OMP-1C protein.

- FIG. 6B shows one embodiment of the OMP-1D protein; FIG. 6A shows one embodiment of the OMP-1D polynucleotide.
- FIG. 7A shows one embodiment of the OMP-1E protein; FIG 7B shows one embodiment of the OMP-1E polynucleotide.
- FIG. 8A shows one embodiment of the OMP-1F protein; FIG 8 B shows one embodiment of the OMP-1F polynucleotide.
- FIG. 9B shows one embodiment of the OMP-1A protein, FIG 9A shows one embodiment of the OMP-1A polynucleotide.
- FIG. 10 B shows one embodiment of a portion of the OMP-1R protein, FIG 10A shows one embodiment of an OMP-1R polynucleotide encoding such polypeptide.
- FIG. 11 B shows one embodiment of a portion of the OMP-1S protein, FIG 11A shows one embodiment of the OMP-1S polynucleotide encoding such polypeptide.
- FIG. 12 B shows one embodiment of a portion of the OMP-1T protein, FIG 12A shows one embodiment of the OMP-1T polynucleotide encoding such polypeptide.
- FIG. 13 B shows one embodiment of the OMP-1U protein, FIG 13A shows one embodiment of the OMP-1U polynucleotide.
- FIG. 14 B shows one embodiment of the OMP-1V protein, FIG 14A shows one embodiment of the OMP-1V polynucleotide.
- FIG. 15 B shows one embodiment of the OMP-1W protein, FIG 15A shows one embodiment of the OMP-1W polynucleotide.
- FIG. 16 B shows one embodiment of the OMP-1X protein, FIG 16A shows one embodiment of the OMP-1W polynucleotide.
- FIG. 17 B shows one embodiment of the OMP-1Y protein, FIG 17A shows one embodiment of the OMP-1Y polynucleotide.
- FIG. 18 B shows one embodiment of the OMP-1Z protein, FIG 18A shows one embodiment of the OMP-1Z polynucleotide.
- FIG. 19 B shows one embodiment of the P30 protein, FIG 19A shows one embodiment of the P30 polynucleotide.
- FIG. 20 B shows one embodiment of the P30a protein, FIG 20A shows one embodiment of the p30A polynucleotide.
- FIG. 21 B shows one embodiment of the P30-1 protein, FIG 21A shows one embodiment of the p30-1 polynucleotide.
- FIG. 22 B shows one embodiment of the P30-2 protein, FIG 22 A shows one embodiment of the p30-2 polynucleotide.

FIG. 23 B shows one embodiment of the P30-3 protein, FIG 23 A shows one embodiment of the p30-3 polynucleotide.

- FIG. 24 B shows one embodiment of the P30-4 protein, FIG 22 A shows one embodiment of the p30-4 polynucleotide.
- FIG. 25 B shows one embodiment of the P30-5 protein, FIG 22 A shows one embodiment of the p30-5 polynucleotide.
- FIG. 26 B shows one embodiment of the P30-6 protein, FIG 26 A shows one embodiment of the p30-6 polynucleotide.
- FIG. 27 B shows one embodiment of the P30-7 protein, FIG 27 A shows one embodiment of the p30-7 polynucleotide.
- FIG. 28 B shows one embodiment of the P30-8 protein, FIG 28 A shows one embodiment of the p30-8 polynucleotide.
- FIG. 29 B shows one embodiment of a portion of the P30-9 protein, FIG 29 A shows one embodiment of the p30-9 polynucleotide encoding such polypeptide.
- FIG. 30 B shows one embodiment of a portion of the P30-10 protein, FIG 30 A shows one embodiment of the p30-10 polynucleotide encoding such polypeptide.
- FIG. 31 depicts the amino acid sequences alignment of seven E. chafeensis OMP-1s and Cowdria ruminantium MAP-1. Aligned positions of identical amino acids with OMP-IF are shown with dots. The sequence of C. ruminantium MAP-1 is from the report of Van Vliet et al (1994) Molecular cloning, sequence analysis, and expression of the gene encoding the immunodominant 32-kilodalton protein of Cowdria ruminantium. Infect. Immun. 62:1451-1456. Gaps indicated by dashes were introduced for optimal alignment of all proteins. Bars indicates semivariable region (SV) and three hypervariable regions (HVI, HV2, and HV3).

DETAILED DESCRIPTION OF THE INVENTION

<u>Isolated Polynucleotides Encoding OMP-1,OMP-1A, OMP-1B, OMP-1C, OMP-1D, OMP-1F and the OMP from E. Canis</u>

In one aspect, the present invention, provides isolated polynucleotides that encode the outer membrane proteins, OMP-1 (or p28), OMP-1B, OMP-1C, OMP-1D, OMP-1E, OMP-1F, OMP-1A, OMP-1R, OMP-1S, OMP-1T, OMP-1U, OMP-1V, OMP-1W, OMP-1X, OMP-1Y and OMP-1Z from E. chafeensis and the outer membrane proteins P30, P30-a, P-30-1, P30-2, P30-3, P30-4, P30-5, P30-6, P30-7, P30-8, P30-9, and P30-10 from E. Canis or an immunogenic fragment thereof.

The polynucleotide is single stranded or double stranded. The polynucleotide may be a DNA or RNA molecule, preferably a DNA molecule, and comprises a sequence which codes for the respective outer membrane protein. Preferably, the polynucleotide encodes at least the mature form of outer membrane protein. The polynucleotide optionally further comprises a leader sequence and encode an outer membrane preprotein that is

processed in the cell to form the mature protein. The polynucleotide of the present invention may also be fused in frame to a marker sequence which allows for purification of the corresponding outer membrane protein.

The OMP-1 polynucleotide encodes an OMP-1 protein of E. chafeensis having a molecular weight of about 27.7 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 3B SEQ ID NO: __; Figure 3B shows one embodiment of the OMP-1 protein, Figure 3A shows one embodiment of the OMP-1 polynucleotide. The OMP-1B polynucleotide encodes an OMP-1B protein of E. chafeensis having a molecular weight of about 28.2 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 4B SEQ ID NO: __; Figure 4B shows one embodiment of the OMP-1B protein, Figure 4A shows one embodiment of the OMP-1B polynucleotide. The OMP-1C polynucleotide encodes an OMP-1C protein of E. chafeensis having a molecular weight of about 27.6 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 5B SEQ ID NO: __; Figure 5B shows one embodiment of the OMP-1C protein, Figure 5A shows one embodiment of the OMP-1C polynucleotide. The OMP-1D polynucleotide encodes an OMP-1D protein of E. chafeensis having a molecular weight of about 28.7 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 6B SEQ ID NO: __; Figure 6B shows one embodiment of the OMP-1D protein, Figure 6A shows one embodiment of the OMP-1D polynucleotide. The OMP-1E polynucleotide encodes an OMP-1E protein of E. chafeensis having a molecular weight of about 27.8 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 7B SEQ ID NO: __; Figure 7B shows one embodiment of the OMP-1E protein, Figure 7A shows one embodiment of the OMP-1E polynucleotide. The OMP-1F polynucleotide encodes an OMP-1F protein of E. chafeensis having a molecular weight of about 27.9 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 8B SEQ ID NO: __; Figure 8B shows one embodiment of the OMP-1F protein, Figure 8A shows one embodiment of the OMP-1F polynucleotide. The OMP-1A polynucleotide encodes an OMP-1A protein of E. chafeensis having a molecular weight of about 29.6 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 9B SEQ ID NO: __; Figure 9B shows one embodiment of the OMP-1A protein, Figure 9A shows one embodiment of the OMP-1A polynucleotide. The OMP-1R polynucleotide encodes an OMP-1R protein of E. chafeensis having a molecular weight of at least 23 kDa and comprising an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 10B SEQ ID NO: __; Figure 10B shows one embodiment of a portion of the OMP-1R protein, Figure 10A shows one embodiment of the OMP-1R polynucleotide encoding such polynucleotide. The OMP-1S polynucleotide encodes an OMP-1S protein of E. chafeensis comprising an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 11B SEQ ID NO: __; Figure 11B shows one embodiment of a portion of the OMP-1S protein, Figure 11A shows one embodiment of the OMP-1S polynucleotide encoding such polypeptide. The OMP-1T polynucleotide encodes an OMP-1T protein of E. chafeensis comprising an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG.12B SEQ ID NO: __; Figure 12B shows one embodiment of a portion of the OMP-1T protein, Figure 12B shows one embodiment of a polynucleotide encoding such polypeptide. The OMP-1U polynucleotide encodes an

OMP-1U protein of E. chafeensis having a molecular weight of about 30.6 kDa and an amino acid sequence which is at least 85% homologous to amino acid sequence shown in FIG. 13B SEQ ID NO: __; Figure 13B shows one embodiment of the OMP-1U protein, Figure 13A shows one embodiment of the OMP-1U polynucleotide. The OMP-1V polynucleotide encodes an OMP-1V protein of E. chafeensis having a molecular weight of about 28.0 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 14B SEQ ID NO: ; Figure 14B shows one embodiment of the OMP-1V protein, Figure 14A shows one embodiment of the OMP-1V polynucleotide. The OMP-1W polynucleotide encodes an OMP-1W protein of E. chafeensis having a molecular weight of about 28.8 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 15B SEQ ID NO: __; Figure 15B shows one embodiment of the OMP-1W protein, Figure 15A shows one embodiment of the OMP-1W polynucleotide. The OMP-1X polynucleotide encodes an OMP-1S protein of E. chafeensis having a molecular weight of about 27.8 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 16B SEQ ID NO: __; Figure 16B shows one embodiment of the OMP-1X protein, Figure 16A shows one embodiment of the OMP-1X polynucleotide. The OMP-1Y polynucleotide encodes an OMP-1Y protein of E. chafeensis having a molecular weight about 28.8 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 17B SEQ ID NO: __; Figure 17B shows one embodiment of the OMP-1Y protein, Figure 17A shows one embodiment of the OMP-1Y polynucleotide. The OMP-1Z polynucleotide encodes an OMP-1Z protein of E. chafeensis comprising an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 18B SEQ ID NO: ; Figure 18B shows one embodiment of a portion of the OMP-1Z protein, Figure 18A shows one embodiment of an OMP-1Z polynucleotide encoding such polypeptide.

The p30 polynucleotide encodes a P30 protein of E. canis having a molecular weight of about 28.8 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 19B SEQ ID NO: Figure 19B shows one embodiment of the P30 protein, Figure 19A shows one embodiment of the p30 polynucleotide. The p30A polynucleotide encodes a P30a protein of E. canis having a molecular weight of about 29.1 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 20B SEQ ID NO: __; Figure 20B shows one embodiment of the P30a protein, Figure 20A shows one embodiment of the p30A polynucleotide. The p30-1 polynucleotide encodes a P30-1 protein of E. canis having a molecular weight of about 27.7 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 21B SEQ ID NO: __; Figure 21B shows one embodiment of the P30-1 protein, Figure 21A shows one embodiment of the p30-1 polynucleotide. The p30-2 polynucleotide encodes a P30-2 protein of E. canis having a molecular weight of about 28.0 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 22B SEQ ID NO: __; Figure 22B shows one embodiment of the P30-2 protein, Figure 22A shows one embodiment of the p30-2 polynucleotide. The p30-3 polynucleotide encodes a P30-3 protein of E. canis having a molecular weight of about 28.7 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 23B SEQ ID NO: __; Figure 23B shows one embodiment of the P30-3 protein, Figure 23A shows one embodiment of the p30-3 polynucleotide. The p30-4 polynucleotide

encodes a P30-4 protein of E. canis having a molecular weight of about 28.0 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 24B SEQ ID NO: __; Figure 24B shows one embodiment of the P30-4 protein, Figure 24A shows one embodiment of the p30-4 polynucleotide. The p30-5 polynucleotide encodes a P30-5 protein of E. canis having a molecular weight of about 29.4 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 25B SEQ ID NO: __; Figure 25B shows one embodiment of the P30-5a protein, Figure 25A shows one embodiment of the p30-5a polynucleotide. The p30-6 polynucleotide encodes a P30-6 protein of E. canis having a molecular weight of about 29.5 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 26B SEQ ID NO: ; Figure 26B shows one embodiment of the P30-6 protein, Figure 26A shows one embodiment of the p30-6 polynucleotide. The p30-7 polynucleotide encodes a P30-7 protein of E. canis having a molecular weight of about 29.9 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 29B SEQ ID NO: __; Figure 29B shows one embodiment of the P30-7 protein, Figure 29A shows one embodiment of the p30-7 polynucleotide. The p30-8 polynucleotide encodes a P30-8 protein of E. canis having a molecular weight of about 30.3 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 28B SEQ ID NO: __; Figure 28B shows one embodiment of the P30-8 protein, Figure 28A shows one embodiment of the p30-8 polynucleotide. The p30-9 polynucleotide encodes a P30-9 protein of E. canis comprising an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 29B SEQ ID NO: __; Figure 29B shows one embodiment of a portion of the P30-9 protein, Figure 29A shows one embodiment of the p30-9 polynucleotide encoding such polypeptide. The p30-10 polynucleotide encodes a P30-10 protein of E. canis comprising an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 30B SEQ ID NO: __; Figure 30B shows one embodiment of a portion of the P30-10 protein, Figure 30A shows one embodiment of the p30-10 polynucleotide encoding such polypeptide.

The polynucleotides encoding an E. chafeensis outer membrane protein or an E. canis outer membrane protein have a sequence that is at least 85%, preferably at least 90%, more preferably at least 95% homologous to or similar to the amino acid sequences shown in Figures 3B through 30B, and thus embrace polynucleotides encoding outer membrane proteins from different strains of E. chafeensis and E. canis. The polynucleotides encode an outer membrane protein whose conserved regions collectively are at least 90%, preferably at 95%, more preferably at least 97% homologous to the conserved regions of the amino acid sequences of the present invention. The outer membrane proteins of E. chafeensis and E. canis have six conserved regions, which are separated by one semivariable region and three hypervariable regions. The conserved regions of the outer membrane proteins OMP-1, OMP-1B, OMP1-C, OMP-1D, OMP1-F are depicted in Fig. 31. Preferably, the amino acid sequence of the outer membrane proteins of E. chafeensis and E. canis are at least 30% divergent from the amino acid sequence of MAP-1. Such sequences include allelic, strain variants and other amino acid sequence variants (e.g., including "muteins" or "mutant proteins"), whether naturally-occurring or biosynthetically produced. As used herein, "amino acid sequence homology" is understood to mean amino acids are conserved amino acids as defined by sequences share identical or similar amino acids, where similar amino acids are conserved amino acids as defined by

Dayoff et al., Atlas of Protein Sequence and Structure; vol. 5, Supp. 3, pp. 345-362 (M. O. Dayoff, ed., Nat'l BioMed. Research Fdn., Washington D.C. 1978.) Thus, a candidate sequence sharing 85% amino acid sequence homology with a reference sequence requires that, following alignment of the candidate sequence with the reference sequence, 85% of the amino acids in the candidate sequence are identical to the corresponding amino acid in the reference sequence, or constitute a conserved amino acid change thereto. "Amino acid sequence identity" is understood to require identical amino acids between two aligned sequences. Thus, a candidate sequence sharing 85% amino acid identity with a reference sequence requires that, following alignment of the candidate sequence with the reference sequence, 85% of the amino acids in the candidate sequence are identical to the corresponding amino acid in the reference sequence.

As used herein, all homologies and identities are calculated using the amino acid sequences shown in the cited Figure or SEQ ID NO as the reference sequence. Thus, to determine whether an amino acid sequence is 85% homologous to OMP-1, one uses the amino acid sequence shown in Fig. ___, SEQ ID NO: ___ as a reference.

Also as used herein, sequences are aligned for homology and identity calculations using the method of the software basic local alignment search tool in the BLAST network service (the National Center for Biotechnology Information, Bethesda, MD) which employs the method of Altschul, S. F., Gish, W., Miller, W., Myers, E. W. & Lipman, D. J. (1990) J. Mol. Biol. 215, 403-410. Identities are calculated by the Align program (DNAstar, Inc.) In all cases, internal gaps and amino acid insertions in the candidate sequence as aligned are ignored when making the homology/identity calculation.

In another aspect, the present invention provides a nucleotide sequence encoding a polypeptide which comprises a fragment of the OMP1 protein, hereinafter referred to as "rP28". The rP28 polypeptide weighs approximately 31 kDa and comprises all but of the first 5 amino acids of mature OMP-1 protein. The rP28 polypeptide comprises the amino acid sequence extending from amino acid 6 through amino acid 251 of the amino acid sequence shown in Fig.1, SEQ ID NO. The present invention also embraces polypeptides where one or more of the amino acids in the sequence extending from amino acid 1 or 6 through amino acid 251 Fig. 1 are replaced by conservative amino acid residues. The present invention also relates to derivatives of rP28 that have an amino acid sequence identity of at least 85%, more preferably at least 90%, and most preferably of at least 95% with the amino acid sequence extending from amino acid 1 or 6 through amino acid 251 of the protein and which derivative binds to antibodies in sera from humans infected with E. chafeensis.

The polynucleotides are useful for producing the outer membrane proteins of E. chafeensis and E. canis. For example, an RNA molecule encoding the outer membrane protein OMP-1 used in a cell-free translation systems to prepare OMP-1. Alternatively, a DNA molecule encoding the outer membrane protein is introduced into an expression vector and used to transform cells. Suitable expression vectors include for example chromosomal, nonchromosomal and synthetic DNA sequences, e.g., derivatives of SV40, bacterial plasmids, phage DNAs; yeast plasmids, vectors derived from combinations of plasmids and phage DNAs, viral DNA such as vaccinia, adenovirus, fowl pox virus, and pseudorabies. The DNA sequence is introduced into the expression vector by conventional procedures.

Accordingly, the present invention also relates to recombinant constructs comprising one or more of the polynucleotide sequences. Suitable constructs include, for example, vectors, such as a plasmid, phagemid, or viral vector, into which a sequence that encodes the outer membrane protein has been inserted. In the expression vector, the DNA sequence which encodes the outer membrane protein is operatively linked to an expression control sequence, i.e., a promoter, which directs mRNA synthesis. Representative examples of such promoters, include the LTR or SV40 promoter, the E. coli lac or trp, the phage lambda PL promoter and other promoters known to control expression of genes in prokaryotic or eukaryotic cells or in viruses. The promoter may also be the natural promoter of the outer membrane protein coding sequence. The expression vector also contains a ribosome binding site for translation initiation and a transcription terminator. Preferably, the recombinant expression vectors also include an origin of replication and a selectable marker, such as for example, the ampicillin resistance gene of E. coli to permit selection of transformed cells, i.e. cells that are expressing the heterologous DNA sequences. The polynucleotide sequence encoding the outer membrane protein is incorporated into the vector in frame with translation initiation and termination sequences. Optionally, the sequence encodes a fusion outer membrane protein which includes an N-terminal or C-terminal peptide or tag that stabilizes or simplifies purification of the expressed recombinant product. Representative examples of such tags include sequences which encode a series of histidine residues, the Herpes simplex glycoprotein D, or glutathione S-transferase.

Polynucleotides which encode portions of the outer membrane proteins of E. chafeensis and E. canus are useful as probes for isolating and identifying E. chafeensis genes and E. canis genes, particularly full-length genes from new strains or isolates of E. chafeensis and E. canis.

The Outer Membrane Proteins of E. chafeensis and E. Canis

In addition to the outer membrane proteins OMP-1, OMP-1B, OMP-1C, OMP-1D, OMP-1 E, and OMP-1F, OMP-1R, OMP-1S, OMP-1T, OMP-1U, OMP-1V, OMP-1W, OMP-1X, OMP-1Y, and OMP-1Z from E. chafeensis and the proteins P30, P30A, P30-1, P30-2, P30-3, P30-4, P30-5, P30-6, P30-7, P30-8, P30-9, and P30-10 from E. Canis, the present inventions embraces non-naturally occurring allelelic forms or derivatives of the outer membrane proteins, where one or more of the amino acids have been replaced by conservative amino acid residues, typically by using direct synthesis or recombinant techniques.

Preparing the Outer Membrane Proteins

The outer membrane proteins of the present invention are synthetically produced by conventional peptide synthesizers. The outer membrane proteins are also produced using cell-free translation systems and RNA molecules derived from DNA constructs that encode the outer membrane protein. Alternatively, the outer membrane protein is made by transfecting host cells with expression vectors that comprise a DNA sequence which encodes the outer membrane protein and then inducing expression of the outer membrane protein in the host cells.

The outer membrane protein is expressed in suitable host cells, preferably bacteria, under the control of suitable promoters. Host cells are transformed with the expression vectors of this invention and cultured in conventional nutrient media. Such media optionally contains additional compounds, such as for example

compounds that induce promoters, such as for example isopropyl-\beta-D-thiogalactoside which induces the Lac promoter, or compounds, such as for example, ampicillin, which allows for selection of transformants.

Following transformation of the suitable host strain and growth of the host strain to an appropriate cell density, the cells are typically harvested by centrifugation, disrupted by physical or chemical means, and the resulting crude extract retained for further purification of the outer membrane protein. Such purification usually involves salting-out of the protein fraction, and one or more chromatography steps, including aqueous ion exchange chromatography, size exclusion chromatography steps, and high performance liquid chromatography (HPLC).

Preparation of Antibodies

The isolated outer membrane proteins, particularly the recombinant forms of the outer membrane proteins, are used as immunogens to produce antibodies immunospecific for the corresponding protein. The term "immunospecific" means the antibodies have substantially greater affinity for the protein used as an immunogen than for other proteins. Such antibodies are generated using conventional techniques by administering the respective outer membrane protein or a portion thereof, i.e., the recombinant polypeptide, to an animal, preferably a nonhuman. collecting blood from the immunized animals and isolating the serum and/or the IgG fraction from the blood. Monoclonal antibodies are prepared by injecting animals with the immunogens, extracting antibody-producing B cells from the animal, fusing the B cells with a myeloma cells to produce hybridomas, obtaining the monoclonal antibodies from the hybridomas.

Antibodies to the outer membrane proteins of E. chafeensis and E. canis are useful research tools for identifying cells, particularly monocytes, infected with E.chafeensis or E. canis and for purifying the corresponding outer membrane protein of E.chafeensis or E. Canis from partially purified preparations by affinity chromatography. Such antibodies are also useful for identifying bacterial colonies, particularly colonies of genetically-engineered bacteria, that are expressing the major outer membrane protein.

Diagnostic Method

The present invention also provides a method for detecting antibodies to the E. chafeensis or E. canis in a sample of a bodily fluid from a patient. The method comprises providing an isolated outer membrane protein of E. chafeensis or E. canis, particularly a recombinant form of the isolated protein, contacting the outer membrane protein or polypeptide with a sample taken from the patient; and assaying for the formation of a complex between the outer membrane protein or polypeptide and antibodies in the sample. For ease of detection, it is preferred that the isolated protein or polypeptide be attached to a substrate such as a column, plastic dish, matrix, or membrane, preferably nitrocellulose. The sample may be a tissue or a biological fluid, including urine, whole blood, or exudate, preferably serum. The sample may be untreated, subjected to precipitation, fractionation, separation, or purification before combining with the isolated protein or peptide. Interactions between antibodies in the sample and the isolated protein or peptide are detected by radiometric, colorimetric, or fluorometric means, size-separation, or precipitation. Preferably, detection of the antibody-outer membrane protein complex is by addition of a secondary antibody that is coupled to a detectable tag, such as for example, an enzyme, fluorophore, or chromophare. Formation of the complex is indicative of the presence of anti-E chafeensis or anti-E canis antibodies,

either IgM or IgG, in the patient. Thus, the method is used to determine whether a patient is infected with E. chafeensis or E. canis.

Preferably, the method employs an enzyme-linked immunosorbent assay (ELISA) or a Western immunoblot procedure. Such methods are relatively simple to perform and do not require special equipment as long as membrane strips are coated with a high quality antigen. Accordingly, it is more advantageous to use a recombinant form of the outer membrane protein of E. chafeensis or E. canis since such proteins, typically, are more pure and consistent in quality than a purified form of such protein.

Immunogenic Composition

The present invention also relates to immunogenic compositions comprising one or more of the isolated outer membrane proteins of E. chafeensis and a pharmaceutically acceptable adjuvant and to immunogenic compositions comprising an isolated P30 protein of E. canis and a pharmaceutically acceptable adjuvant, which, preferably, enhances the immunogenic activity of the outer membrane protein in the host animal.

Preparation of a Polynucleotide which Encodes OMP-1(P28)

A. Isolation of the Outer Membrane Proteins

E. chafeensis Arkansas strain and E. canis Oklahoma strain were cultivated in the DH82 dog macrophage cell line and purified by Percoll density gradient centrifugation. Purified ehrlichiae (100 μg) were suspended with 10 mM sodium phosphate buffer, pH 7.4, containing 0.1% Sodium N-lauroyl sarcosine (Sarkosyl) [Sigma, St. Louis, MO], 50 μg/ml each Dnase I (Sigma) and Rnase A (Sigma), and 2.5 mM MgCl₂. After incubation at 37° for 30 min, the sample was separated by centrifugation at 10,000 x g for 1 h into the soluble supernatant and the insoluble precipitate. The insoluble pellet was resuspended 2 to 3 times with 0.1% Sarkosyl and centrifuged. The final pellet was analyzed by sodium dodecyl sulfate (SDS)-polyacrylamide gel electrophoresis (PAGE) and by electron microscopy.

Transmission electron microscopy revealed that the purified ehrlichial fraction consists of a mixture of electron dense and light forms of *E. chafeensis* with slight disintegration of inner membrane. Ehrlichiae were not surrounded with the host inclusion membrane. Various sizes of membrane vesicles (< 1 µm) without significant ribosomes or nuclear materials were observed in the Sarkosyl-insoluble fraction from the organism. Succinic dehydrogenase (inner membrane marker enzyme of gram negative bacteria) activities were at less than the detection limit (1 n moles / min / mg of protein) in the Sarkosyl-insoluble fraction compared to approximately 10 n moles / min / mg of protein in the Percoll-purified organisms, suggesting that the insoluble fraction primarily consisted of the outer membrane of *E. chafeensis*.

Analysis of the Sarkosyl-soluble, and insoluble fraction of *E. chafeensis* by SDS-PAGE suggested that proteins of 30-kDa range in the insoluble fraction represent the major outer membrane proteins of this organism. Analysis of the Sarkosyl-soluble, and insoluble fraction of *E. canis* by SDS-PAGE suggested that proteins of 30-kDa range in the insoluble fraction represent the major outer membrane proteins of this organism also. *E. canis* was

antigenically cross reactive with *E. chafeensis*. These findings indicate that the 30-kDa range proteins represent the major outer membrane proteins of these two *Ehrlichia* spp.

To improve resolution of the outer membrane proteins, proteins in the Sarkosyl-insoluble pellet prepared from 400 µg of purified *E. chafeensis* were separated by a reversed-discontinuous (Rd) SDS-PAGE (2.5-cm-long 17% gel on top of 11-cm-long 12% gel). At least five proteins of 30-kDa range in *E. chafeensis* (P23, P25, P27, P28, and P29) were resolved from the Sarkosyl-insoluble proteins.

B. Cloning and sequencing of the p28 gene

The	portion of the	nembrane contain	ing bound prot	eins was exc	ised and analyzed	with a	n App	oned
Biosystems pr	rotein sequencer	(Model 470). The	N-terminal ami	no acid seque	nce of P28 was de	termined	i as D	P A
GSGING	NFYSGKY	M P, SEQ IN NO	D Ba	sed on 6th to	12th amino acids	of this s	equen	ce, a
forward	primer,	FECH1,	having -	the	sequence:			5'-
CGGGATCC	GAATTCGG(A/	T/G/C)AT(A/T/C)	AA(T/C)GG(A/	T/G/C)AA(T/	C)TT(T/C)TA-3'.	SEQ	ID	NO
was	designed. Ami	no acids at the 1	o 5 positions o	f the N termin	nus of P28 were n	ot inclu	ded in	ı this
primer design	n. For insertion	into an expression	vector, a 14-bp	sequence (ur	iderlined) was add	ed at the	: 5' er	nd of
primer to crea	ate an <i>Eco</i> RI and	l a <i>Bam</i> HI site. Th	e reverse prime	, RECH2, w	hich includes a No	il site at	the 5	' end
for ligation in	nto an expression	n vector had the s	equence: 5'-A	GCGGCCGC	TTA(A/G)AA(T/C)A(C/G) (A/C	3)AA
(C/T)CT T(C	/G)C TCC-3'. S	EQ ID NO	•					

Genomic DNA of *E. chafeensis* was isolated from purified organisms. PCR amplification with FECH1 and RECH2 primers was performed using a Perkin-Elmer Cetus DNA Thermal Cycler (model 480). A 0.8-kb amplified product was cloned in the pCRII vector of a TA closing kit, as described by the manufacturer (Invitrogen Co., San Diego, CA). The clone obtained was designated pCRIIp28. Both strands of the inserted DNA were sequenced by a dideoxy-termination method with an Applied Biosystems 373A DNA sequencer.

The 0.8-kb DNA fragment, cloned in pCRIIp28, had an open reading frame (ORF) of 756 bp encoding a 251-amino acid recombinant protein (including both PCR primer regions) with a molecular mass of 27,685 Da. The nucleotide sequence of the open reading frame, SEQ ID NO: , and the amino acid sequence of the polypeptide of the OMP-1 protein, SEQ ID NO ___, are shown in Figs ____ and _____, respectively.

A DNA fragment comprising the p30 gene was prepared in a similar manner, i.e., by PCR amplification of genomic DNA of E. canis with the FECH1 and RECH2 primers.

Preparation of Polynucleotides which encode OMP 1A, OMP1B, OMP1-C, OMP-1D, OMP-1F, and OMP1-E

A. Southern blot analysis. Genomic DNA extracted from the purified *E. chafeensis* (200 ng each) was digested with restriction endonucleases, electrophoresed, and transferred to Hybond-N⁺ nylon membrane (Amersham, Arlington Heights, IL), by a standard method. The 0.8-kb *p28* gene fragment from the clone pCRII*p28* was labeled with [α-³²P]dATP by the random primer method using a kit (Boehringer Mannheim, Indianapolis, IN) and the labeled fragment was used as a DNA probe. Hybridization was performed at 60°C in rapid hybridization buffer (Amersham) for 20 h. The nylon sheet was washed in 0.1 x SSC (1 x SSC containing 0.15M sodium chloride and

0.015M sodium citrate) with 1% SDS at 55°C and the hybridized probes were exposed to Hyperfilm (Amersham) at -80°C.

Genomic Southern blot analysis with several restriction enzymes resulted in one or more DNA fragment(s) of *E. chafeensis* which hybridized to ¹²P-labeled *p28* gene probe. The restriction enzymes used did not cut within the *p28* gene portion of the pCRII*p28* insert. *Xba* I, *BgI* II, and *Kpn* I produced two bands, *Spe* I generated three bands, and *EcoR* V and *Pst* I produced multiple bands with different densities. *EcoR* I generated a broad band of 2.5 to 4kb. These *p28* homologous genes are designated as *omp-1* (outer membrane protein-1) family.

B. Cloning and sequencing of genomic copies of *E. chafeensis p28* gene. The *EcoR* I and *Pst* I fragments of DNA, detected by genomic Southern blot analysis as described above, were inserted into pBluescript II KS (+) vectors, and the recombinant plasmids were introduced into *E. coli* DH5α. Using the colony hybridization method with the ³²P-labeled *p28* gene probe, four positive clones were isolated from the transformant. The positive clones were designated pEC2.6, pEC3.6, pPS2.6, and pPS3.6. These contained the ehrlichial DNA fragments of 2.6-kb (*EcoR* I), 3.6 kb (*EcoR* I), 2.6 kb (*Pst* I), and 3.6 kb (*Pst* I), respectively. The inserts of the clones pEC3.6 and pPS2.6 overlapped as shown in Fig. _____. The overlapping area was further confirmed by PCR of *E. chafeensis* genomic DNA with two pairs of primer sets interposing the junctions of the four clones. The 1.1- to 1.6-kb DNA fragments of *Hind*III-*Hind*III, *Hind*III-*EcoR*I, or *Xho*I-*EcoR*I in the pEC2.6 and pEC3.6 were subcloned for sequencing. DNA sequencing was performed with suitable synthetic primers by dideoxy-termination method as described above.

Four DNA fragments from 2.6 to 3.6 kb were cloned from the *Eco*RI-digested and the *Pst*I-digested genomic DNA of *E. chafeensis* by colony hybridization with radiolabeled *p28* gene probe. The inserted DNA of the two recombinant clones, pEC3.6 and PPS2.6, were overlapped as shown in Fig. 7. Sequencing revealed one 5'-truncated ORF of 243 bp (designated *omp-1A*) and five complete ORF of 836-861 bp (designated *omp-1B* to *omp-1F*), which are tandemly-arrayed and are homologous to the *p28* gene (but are not identical), in the ehrlichial genomic DNA of 6,292 bp. The intergenic spaces were 581 bp between *omp-1A* and *omp-1B* and 260-308 bp among others. Putative promoter regions and ribosome-binding sites were identified in the noncoding regions.

Sequence analysis and GenBank accession number.

Nucleotide sequences were analyzed with the DNASIS program (Hitachi Software Engineering Co., Ltd., Yokohama, Japan). A homology search was carried out with databases of the GenBank, Swiss Plot, PDB and PIR by using the software basic local alignment search tool in the BLAST network service (the National Center for Biotechnology Information, Bethesda, MD). Phylogenetic analysis was performed by using the PHYLIP software package (version 3.5). An evolutional distance matrix, generated by using the Kimura formula in the PROTDIST, was used for construction of a phylogenetic tree by using the unweighted pair-group method analysis (UPGMA) (Felsenstein, J. 1989. PHYLIP-phylogeny inference package (version 3.3). Cladistics 5:164-166). The data were also examined using parsimony analysis (PROTPARS in PHYLIP). A bootstrap analysis was carried out to investigate the stability of randomly generated trees by using SEQBOOT and CONSENSE in the same package. The nucleotide sequence of the *p28* gene and its gene copies has been assigned GenBank accession numbers U72291 and AF021338, respectively.

Proteins of the E. chafeensis omp-1 Family.

Five complete *omp-1* gene copies (*omp-1B* to *omp-1F*) encode 279 to 287-amino acid proteins with molecular masses of 30,320 - 31,508 Da. *Omp-1A* encodes an 82-amino acid partial protein (9,243 Da) which lacks the N-terminal region. The 25-amino acid sequence at the N-terminus of OMP-1B to OMP-1F (encoded in *omp-1B* to *omp-1F*) is predicted to be a signal peptide because three carboxyl-terminal amino acids of the signal peptides (Ser-X-Ala in OMP-1B, Leu-X-Ser for OMP-C, and Ser-X-Ser for OMP-1D and OMP-1F) are included in the preferred amino acid sequence of signal peptidase at the processing sites proposed by Oliver .. The calculated molecular masses of the mature OMP-1B to OMP-1F from the predicted amino acid sequences are 28,181 Da for OMP-1B, 27,581 Da for OMP-1C, 28,747 Da for OMP-1D, 27,776 Da for OMP-1E, and 27,933 Da for OMP-1F. The estimated isoelectric points are 4.76-5.76 in the mature OMP-1B to OMP-1F. An amino acid sequence in *omp-1F* gene (the 80th to 94th amino acids) was identical to the N-terminal amino acid sequences of *E. chafeensis* native P23 protein as determined chemically, which indicates that P23 is derived from the *omp-1F* gene. Amino acid sequences identical to the N-terminal sequences of P25, P27, and P29 were not found in those from *omp-1* gene copies cloned in this study.

Alignment of predicted amino acid sequences of the *E. chafeensis* OMP-1 family and *Cowdria ruminantium*, revealed substitutions or deletions of one or several contiguous amino acid residues throughout the molecules. The significant differences in sequences among the aligned proteins are seen in the regions indicated SV (semivariable region) and HV (hypervariable region) 1 to 3 in Fig 31. Computer analysis for hydropathy revealed that protein molecules predicted from all *omp-1* gene copies contain alternative hydrophilic and hydrophobic motifs which are characteristic of transmembrane proteins. The HV1 and HV2 were found to locate in the hydrophilic regions.

The amino acid sequences of 5 mature proteins without signal peptides (OMP-1C to OMP-1F and a P28) were similar to one another (71-83%) but the sequence of OMP-1B was dissimilar to those of the 5 proteins (45-48%). The amino acid sequences of the 5 proteins showed an intermediate degree of similarity with that of C. ruminantium MAP-1 (59-63%), but the similarity between that of the OMP-1B and the C. ruminantium MAP-1 was low (45%). These relations are shown in a phylogenetic tree which was obtained based on the amino acid sequence alignment by UPGMA method in the PHYLIP software package (Fig. 10). Three proteins (P28, OMP-1D, and OMP-1F) and two proteins (OMP-1C and OMP-1E) formed two separate clusters. The OMP-1B was located distantly from these two clusters. The C. ruminantium MAP-1 was positioned between the OMP-1B and other members in the OMP-1 family.

Preparation of a Recombinant form of OMP-1 and P30

The 0.8-kb p28 gene was excised from the clone pCRIIp28 by EcoRI-NotI double-digestion, ligated into EcoRI-NotI sites of a pET 29a expression vector, and amplified in Escherichia coli BL21 (DE3)pLysS (Novagen, Inc., Madison, WI). The clone (designated pET29p28) produced a fusion protein with a 35-amino acid sequence

carried from the vector at the N terminus. The amino acid sequence of the OMP-1 portion of the fusion protein is depicted in Fig. 1.

An expression vector comprising the p30 gene was used to prepare the recombinant form of P30.

The following examples are for purposes of illustration only and are not intended to limit the scope of the claims which are appended hereto.

Preparation of anti rP28 (anti-OMP1) antibody

The (r) P28 antigen was prepared by excising the gel band corresponding to the rP28 in SDS-PAGE, mincing the band in phosphate-buffered saline (PBS), pH 7.4, and mixing with an equal volume of Freund's incomplete adjuvant (Sigma). The rP28 mixture (1 mg of protein each time) was subcutaneously injected into a rabbit every 2 weeks four times. A serum sample was collected from the rabbit to provide the anti-rP28 antibody

The anti-rP28 antibody was examined by western immunoblots analysis. The results indicated that the rabbit anti-rP28 antibody recognized not only rP28 (31 kDa) and P28, but also P29 and P25 of E. chafeensis and P30 of E. canis. These results indicate that P28 shares antigenic epitopes with P25 and P29 in E. chafeensis and P30 of E. canis.

Example 1. Assaying for the presence of anti-OMP-1 antibody in a Patient

Convalescent-phase serum from a patient with clinical signs of human ehrlichiosis was used. Western blot analyses using the rP28 protein as antigen was performed with 1:1,000 dilutions of this serum. Alkaline phosphatase-conjugated affinity-purified anti-human, anti-rabbit or anti-mouse immunoglobulin G (Kirkegaard & Perry Laboratories, Inc., Gaithersburg, MD) were used at a 1:1,000 or 1:2,000 dilution as secondary antibodies. Results indicated that serum from a patient with clinical signs of human ehrlichiosis reacted strongly to rP28 (31 kDa).

Example 2 Assaying for the presence of anti-OMP-1 antibody in a Patient

Convalescent-phase serum from a patient with clinical signs of human ehrlichiosis was reacted with the rP30 protein of E.canis as described in Example 1. The serum reacted strongly to rP30. These results indicate the rP30 is useful for diagnosing an infection with E. chafeensis in human patients.

Example 3. Identifying E. chafeensis-infected cells using anti-rP 28 antibody

E. chafeensis-infected DH82 cells were sonicated and centrifuged at 400 x g for 10 min. The supernatant was then centrifuged at 10,000 x g for 10 min to obtain ehrlichia-enriched pellet. The pellet was resuspended and incubated with rabbit anti-rP28 antibody or normal rabbit serum (1:100 dilution) at 37°C for 1h in PBS containing 1% bovine serum albumin (BSA-PBS). After washing, the ehrlichiae was incubated with gold-conjugated protein G (20 nm), Sigma) at 1:30 dilution for 1 h at room temperature in BSA-PBS. After washing again, the specimen was fixed with 1.25% formaldehyde, 2.5% glutaraldehyde, and 0.03% trinitrophenol in 0.1 M cacodylate buffer (pH 7.4) for 24h and postfixed in 1% osmium-1.5% potassium ferricyanide for 1 h (34). The section was then embedded in

PolyBed 812 (Polysciences, Warraington, Pa). The specimen was ultrathin sectioned at 60 nm, stained with uranyl acetate and lead citrate, and observed with a Philips 300 transmission electron microscope at 60 kV.

Transmission immunoelectron microscopy with colloidal gold-conjugated protein G and rabbit anti-rP28 antibody revealed gold particles bound to *E. chafeensis* surface. The distribution of the particles was random, close to the surface, and appeared as if almost embedded in the membrane, suggesting that the antigenic epitope protrudes very little from the lipid bilayer. Nonetheless, the antigenic epitope was surface-exposed, and thus, could be recognized by rabbit anti-rP28 antibody. No gold particles were observed on host cytoplasmic membrane or *E. chafeensis* incubated with normal rabbit serum.

Example 4. Immunization of mice and E. chafeensis challenge.

4).

The rP28 band in SDS- PAGE was excised, minced, and mixed with an equal volume of Freund's incomplete or complete adjuvant. Nine BALB/c male mice (6 weeks old) were divided into two groups. Five mice were intraperitoneally immunized a total of four times at 10-day intervals; twice with a mixture of the minced gel with the rP28 (30 to 40 µg of protein per mouse each time) and incomplete adjuvant, and twice with a mixture of the recombinant protein (the same amount as before) and complete adjuvant. Four mice were intraperitoneally injected with a mixture of the minced gel without protein and the respective adjuvants. For ehrlichia-challenge, approximately 1 x 10⁷ DH82 cells heavily-infected with *E. chafeensis* were disrupted by sonication in serum-free DMEM (GIBCO-BRL) and centrifuged at 200 x g for 5 min. The supernatant was diluted to a final volume of 5 ml, and 0.3 ml was inoculated intraperitoneally into each mouse 10 days after the last immunization. Before challenge, all 5-immunized mice had a titer of 1:160 against *E. chafeensis* antigen by IFA and all 4-nonimmunized mice were negative.

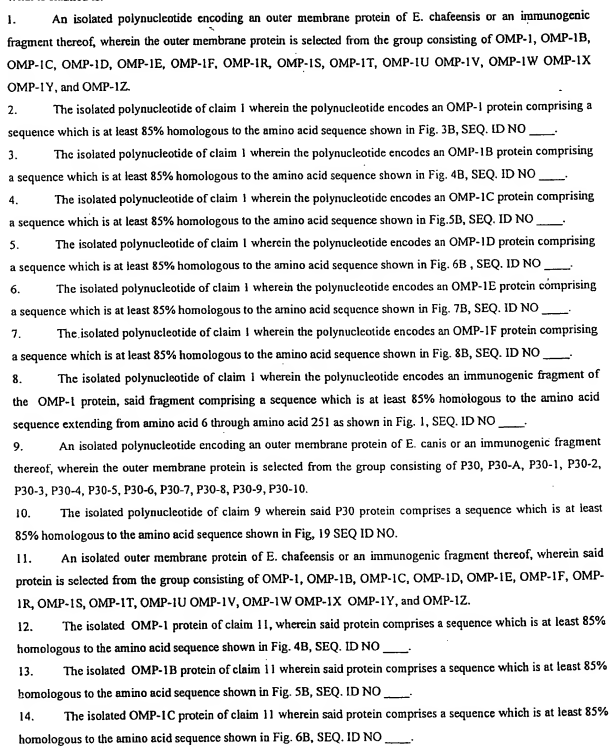
At day 5 post-challenge, approximately 1 ml of blood was collected in an EDTA tube from each mouse and protection was assessed by PCR detection of *E. chafeensis* 16S rDNA in the buffy coat of the collected blood. *E. chafeensis* could not be reisolated in cell culture at day 10 postinfection. Day 5 post challenge is the optimum time at which establishment of ehrlichial infection can be examined by PCR without the influence of residual DNA from the ehrlichiae used as the challenge before the spontaneous clearance of organisms take place. The *E. chafeensis*-specific DNA fragment was observed in all nonimmunized mice but not in any immunized mice, indicating that immunization of rP28 apparently protects mice from ehrlichial infection and indicating that the P28 is a potential protective antigen

Example 5 Assaying for the presence of anti-P30 antibody in Dogs

The rP30 protein was used as an antigen in a Western immunoblot analysis and dot blot analysis to detect the presence of antibody to E. canis in serum from E-canis infected dogs. The results of the Western immunoblot analysis indicated that reactivity of the sera with rP30 was stronger than the reactivity that was observed when purified E.canis was used as antigen. The results of the dot blot assay indicated that rP30 is a useful and sensitive tool for serodiagnosis of canine ehrlichiosis.

CLAIMS

What is claimed is:



The isolated OMP-1D protein of claim 11 wherein said protein comprises a sequence which is at least 85% 15. homologous to the amino acid sequence shown in Fig. 7B, SEQ. ID NO ___ The isolated OMP-1E protein of claim 11 wherein said protein comprises a sequence which is at least 85% 16. homologous to the amino acid sequence shown in Fig. 8B, SEQ. ID NO ____. The isolated OMP-1F protein of claim 11 wherein said protein comprises a sequence which is at least 85% 17. homologous to the amino acid sequence shown in Fig. 9B, SEQ. ID NO ____. The isolated immunogenic fragment of the OMP-1 protein of claim 11, said fragment comprising a sequence which is at least 85% homologous to the amino acid sequence extending from amino acid 6 through amino acid 251 as shown in Fig. 1, SEQ. ID NO ____. An isolated outer membrane protein of E. canis or an immunogenic fragment thereof, wherein the outer 19. membrane protein is selected from the group consisting of P30, P30-A, P30-1, P30-2, P30-3, P30-4, P30-5, P30-6, P30-7, P30-8, P30-9, P30-10. The isolated P-30 protein of claim 19 wherein said protein comprises a sequence which is at least 85% 20. homologous to the amino acid sequence shown in Fig 19, SEQ ID NO. . . A method for diagnosing an infection with E. chafeensis in a patient comprising the steps of: 21. (a) providing a serum sample from the patient; of claim (b) providing an outer membrane protein selected from the group consisting of a protein 11, a protein of claim 19, and mixtures thereof; (c) contacting the serum sample with the outer membrane protein; and (d) assaying for the formation of a complex between antibodies in the serum sample and the outer membrane protein, wherein formation of said complex is indicative of infection with E. chafeensis. A method for diagnosing an infection with E. canis in a Canidae patient comprising the steps of: 22. (a) providing a serum sample from the patient; (b) providing an outer membrane protein of claim 19; (c) contacting the serum sample with the outer membrane protein; and (d) assaying for the formation of a complex between antibodies in the serum sample and the outer membrane protein, wherein formation of said complex is

indicative of infection with E. canis.

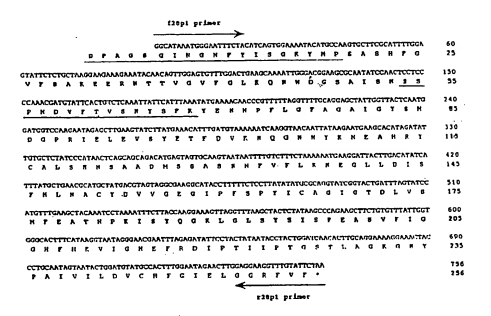


Fig. 1

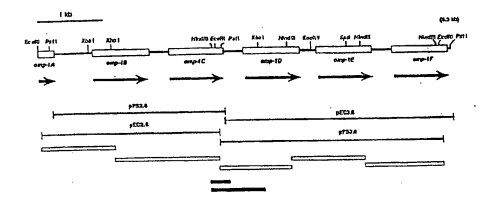


Fig. 2

				20	10
	50	40	30	AAAAAGTTTT	
TTCTCTACCT	CATTAATATC				70
. 120	110	100	90	80	GGAGTATCAT
CATCAGTGGA	GTAATTTCTA		AGCAGGTAGT	TTTCCGACCC	130
. 180	· 170	160	150	140	
AAGAAATACA	CTAAGGAAGA	GTATTCTCTG	GCATTTTGGA	CAAGTGCTTC	190
240	230	220	210	200	ACAGTTGGAG
CAACTCCTCC	GCGCAATATC		GAAGCAAAAT		T
300	290	280	270	260	250
	ATGAAAACAA	TCATTTAAAT	CTCAAATTAT		CCAAACGATG
360	350	. 340	330	320	310
TGAAGTATCT	GAATAGAGCT	GATGGTCCAA	TTACTCAATG		GGTTTTGCAG
420	410	400	390	380	370
ACATAGATAT	AGAATGAAGC	AACAATTATA	AAATCAAGGT	TTGATGTAAA	TATGAAACAT
.400	470	460	450	440	430
TTTTCTCTTT	CAAGTAATAA	ATGAGTAGTG	AGCAGCAGAC	CCCATAACTC	TGTGCTCTAT
540	530	520	510	500	490
TGACGTAGTA	ACGCATGCTA	TTTATGCTGA	TGACATATCA	AAGGATTACT	
	590	580	570	560	550
TTTAGTATCC	TCGGTACTGA	TGCGCAGGTA	TCCTTATATA	TACCTTTTTC	GGCGAAGGCA
660	650	640	630	620	610
AAGCTACTCT		TACCAAGGAA	TAAAATTTCT	CTACAAATCC	ATGTTTGAAG
720	710	700	690	680	670
AGGGAACGAA	ATAAGGTAAT	GGGCACTTTC	GTTTATTGGT	AAGCTTCTGT	ATAAGCCCAG
780	770	760	750	740	730
	TTGCAGGAAA	GGATCAACAC	AATACCTACT	TTCCTACTAT	TTTAGAGATA
840	830	820	810	800	790
A A C C T T T C T A	AACTTGGAGG	TTTGGAATAG	TGTATGCCAC	TAATACTGGA	CCTGCAATAG
900	890	880	870	860	850
900			•••••	• • • • • • • • • • • • • • • • • • • •	TTCTAA
	• • • • • • • • •	·			

Fig. 3A

. 10	20	30	40	50	60
MNYKKVFITS	ALISLISSLP	GVSFSDPAGS	GINGNFYISG	KYMPSASHFG	VESAKEERNT
70	80	90	100	110	120
TVGVFGLKQN	WDGSAISNSS	PNDVFTVSNY	SFKYENNPFL	GFAGAIGYSM	DGPRIELEVS
130	140	150	160	. 170	180
YETFDVKNQG	NNYKNEAHRY	CALSHNSAAD	MSSASNNEVE	LKNEGLLDIS	FMLNACYDVV
190	200	210	220	230	240
GEGIPFSPYI	CAGIGTDLVS	MFEATNPRIS	YQGKLGLSYS	ISPEASVFIG	GHFHKVIGNE
250	260	270	280	290	300
FRDIPTIIPT	GSTLAGKGNY	PAIVILDVCH	FGIEL-GGREV	F	••••

Fig. 3B

10	20	30	40	50	60
ATGAATTACA	AGAAAATTTT	TGTAAGCAGT	GCATTAATTT	CATTAATGTC	
70	80	90	100	110	120
TACCAATCTT	TTGCAGATCC	TGTAACTTCA	AATGATACAG	GAATCAACGA	
130	140	150	160	170	. 180
GGCTTCTACA	TTAGTGTAAA	GTATAATCCA	AGCATATCAC	ACTTCAGAAA	ATTCTCAGCT
190	200	210	220	230	240
GAAGAAGCTC	CCATCAATGG	AAATACTTCT	ATCACTAAAA	AGGTTTTCGG	GCTGAAAAA
250	260	270	280	290	300
GACGGAGATA	TAGCACAATC	TGCGAATTTT	AACAGGACAG	ATCCAGCCCT	CGAGTTTCAG
310	320	330	340	350	360
AATAACCTAA	TATCAGGATT	CTCAGGAAGT	ATTGGTTATG	CTATGGATGG	GCCAAGAATA
370	380	390	400	410	420
GAACTTGAAG	CTGCATACCA	AAAATTTGAT	GCAAAAAATC	CTGACAACAA	TGACACTAAT
430	440	450	460	470	480
AGCGGTGACT	ACTATAAATA	CTTTGGACTA	TCTCGTGAAG	ACGCAATAGC	AGATAAGAAA
490	500	510	520	530	.540
TATGTTGTCC	TTAAAAATGA	AGGCATCACT	TTTATGTCAT	TAATGGTTAA	CACTTGCTAT
550	560	570	-580	590	600
GACATTACAG	CTGAAGGAGT	ACCTTTCATA	CCGTATGCAT	GTGCAGGTGT	AGGAGCAGAC
610	620	630	640	650	. 660
CTTATAAACG	TATTTAAGGA	TTTTAATTTA	AAATTCTCAT	ACCAAGGGAA	AATAGGTATT
670	680	690	700	710	720
AGCTATCCA	TCACACCAGA	AGTTTCCGCT	TTTATTGGA	GATACTACCA	CGGAGTTATA
730					
GGAAATAAT	TTAACAAAAT	ACCTGTAATA	ACACCTGTA	TATTAGAAG	AGCTCCTCAA
790				•	
ACCACATCT	G CGCTAGTAAC	TATTGACACT	GGATACTTT	G GCGGAGAAG!	r TGGAGTAAGG
850	860	870	88	0 89	900
TTCACCTTC'	r AG				
	•	Fig	g. 4A		
					n 60
10				-	•
				E GFYISVKYN	P SISHFRKESA
70					
					S IGYAMDGPRI 0 180
130				0 17	•
					T FMSLMVNTCY
19					0 240.
					A FIGGYYHGVI
25				0 29	
GNNFNKIPV	I TPVVLEGAP	Q TTSALVTID	T GYFGGEVGV	R FTF	

Fig. 4B

10	20		40	50	60
					TTTCTTACCT
70	BO BO	90	100	110	120
GGAATATTAC	TTTCTGAACC		GACAGTGTGA		CTATATTAGT
130	140	AGIACAAGAT	160	170	180
	TGCCAAGTGC			CTGCCAAAGA	,
GGCAAGTACA 190	200	210	220	230	240
CCTACTGTCG	CGTTGTATGG	,			TTCAAGTCAT
250	260	270	280	290	300
GCTGATGCGG		CAAAGGTTAT		ACGAAAACAA	
310	320	330	340	350	360
GGTTTTGCAG				GAATAGAGTT	
370	380	390	400	410	420
TATGAAACAT	• • • • • • • • • • • • • • • • • • • •			AAAATGATGC	
430	440	450	460	470	480
TGTGCCTTAG				CTAGTCACTA	
490	500	510	520	530	540
ÄAAAATGAAG				-	CGTAGTAAGT
550	560	570	580		600
GAAGGAATAC					-
610		630			•
TTTGAAGCTA		. AATTTCTTAT			
670					:-
AACCCAGAAG		-		AAGTTGCAGG	7
730					
AGGGACATTI	CTACTCTTAA	AGCGTTTGCT		CTGCAGCTAC	TCCAGACTTA
790	800	810	82	830	840
GCAACAGTA	CACTGAGTGT	GTGTCACTT	GGAGTAGAA	C TTGGAGGAA	ATTTAACTTC
850	860	870) 88	0 890	900
TAA			• • • • • • • • •		

Fig. 5A

60	50	40	30	20	. 10
GVFSAKEEKN	GKYMPSASHF	DSVSGNFYIS	GILLSEPVQD	ALALPMSFLP	MNCKKFFITT
120	110	100	90	. 80	70
GGPRIEFEVS	GFAGAIGYSM	SFKYENNPFL	ADADFNNKGY	DWNGVSASSH	PTVALYGLKQ
180	170	. 160	150	140	130
MLNACYDVVS	KNEGLLDISL	NATASHYVLL	CALDRKASST	GNYKNDAHRY	YETFDVKNQG
240	230	220	210	. 200	190
HFHKVAGNEF	NPEASVFVGG.	QGKLGLSYSI	FEAINPKISY	AGVGTDLISM	EGIPFSPYIC
300	290	280	270	260	250
		GVELGGRENE	ATVTLSVCHF	TPSSAATPDL	RDISTLKAFA

Fig. 5B

10	20	30	40	50	
ATGAACTGCG	AAAAATTTTT				60 CTTCTTACCT
70	. 80	90	100		
GGAATATCAC	TTTCTGATCC	AGTACAGGAT	TOU	110	120
130	140	150	160		CTACATCAGT
GGAAAGTATA	TGCCAAGCGC	TTCGCATTTT	டுத்தோரையாம	170	180
190	200	210	220		
ACAACAGTTG	GAGTATTTGG	AATAGAGCAA	CATTCCCNTN	230	240
250	260	270	280		ATCTAGAACC
ACTTTAAGCG	ATATATTCAC	CGTTCCAAAT	200 ጥልጥጥድ ግጥጥጽ	290	300
310	320	330	340		
TCAGGATTTG	CAGGAGCTAT	TGGCTACTCA	ATCCATCCCC	350	360
370	380	390	400		
TCTTATGAAG	CATTCGATGT	TAAAAATCAA	CCTT A CA A MM	410	420
430	440	450	460		
TATTATGCTC		TCTCGGCACA	UOP CDCDCDCDCD	470	480
490	500	510	520		
TCTGTCTTTC	TAATAAATGA		CAMARAMORM	530	540
550	560	570	580		
GATGTAATAA	GTGAAGGCAT	ACCTTTTTTTCT		590	600
610	620	630	640		
TTAGTATCCA	TGTTTGAAGC		D4U	650	660
670	680	690	700		
AGTTACCCTA	TAAGCCCAGA		יייייי איייייייייייייייייייייייייייייי	710	720
730	740	750	760		
GGAAACGAAT	TTAGAGATAT		UO /	770	780
790	800	810			
GGAAACTACC	CTGCAATAGT	AACACTGGAG	820	830	840
850	860	870	GIGITCTACT		
AGGTTTAACT	TCCAACTTTG	A	880	890	900
			• • • • • • • • • •	• • • • • • • • •	

Fig. 6A

10	20	30	40	50	60
MNCEKFFITT	ALTLLMSFLP	GISLSDPVQD	DNISGNEYIS	GKYMPSASHF	GVFSAKEERN
70	80	90	100	110	120
rtvgvfgieq	DWDRCVISRT	TLSDIFTVPN	YSFKYENNLF	SGFAGAIGYS	MDGPRIELEV
130	140	150	160	170	180
Syeafdyknq	GNNYKNEAHR	YYALSHLLGT	ETQIDGAGSA	SVFLINEGLL	DKSFMLNACY
190	. 200	210	220	230	240
DVXSEGIPFS	PYICAGIGID	LVSMFEAINP	KISYQGKLGL	SYPISPEASV	FIGGHFHKVI
250	260	270	280	290	300
GNEFRDIPTM	IPSĖSALAGK	GNYPAIVTLD	VFYFGIELGG	RENEOL	

Fig. 6B

10	20	30	40	50	60
ATGAATTGCA	AAAAATTTTT	TATAACAACT	GCATTAGTAT	CACTAATGTC	CTTTCTACC
70	б́в	90	100	110	120
GGAATATCAT	TTTCTGATCC	AGTGCAAGGT		GTGGTAATTT	CTATGTTAGT
130	140	150	160	170	180
GGCAAGTATA	TGCCAAGTGC	TTCGCATTTT		CTGCCAAAGA	
190	200	210	220	230	240
CCTACTGTTG	CATTGTATGG	CTTAAAACAA		GGATTAGCTC	ATCA ACTOR
250	260	270	280	290	300
AATGATAATC	ATTTCAATAA	CAAGGGTTAT		ATGAAAATAA	
310	320	330	340	350	360
GGGTTTGCAG	GAGCTATTGG	TTATTCAATG			TGAAGTGTCC
370	380	390	400	410	
TATGAAACAT	TTGACGTTAA	AAATCAGGGT			420 TCACAGATAC
430	440	450	460	47.0	
TGTGCTTTAG	GTCAACAAGA	CAACAGCGGA		CTAGTAAATA	480
490	500	510	520	530	
AAAAGCGAAG	GATTGCTTGA			CATGCTATGA	540
550	560	570	580	590	
GAGAGCATAC	CTTTGTCTCC	· · · · -			AATATCCATG
610	620	630	640	650	660
TTTGAAGCTA	CAAATCCTAA	AATTTCTTAC		TAGGTCTAAG	TTACTCTATA
670	680	690	700	710	720
AACCCAGAAG	CTTCTGTATT	TATTGGTGGA			AAACGAATTT
730	740	750	760	770	780
AGGGACATTC	CTACTCTGAA	AGCATTTGTT		CTACTCCAGA	
790	800	810	820	830	840
GTAACACTAA	GTGTATGTCA		GAACTTGGAG	GAAGGTTTAA	СТТСТА

Fig. 7A

. 10	20	30	40	50	. 60
MNCKKFFITT	ALVSLMSFLP	GISFSDPVQG	DNISGNEYVS	GKYMPSASHF	GMESAKEEKN
70	90	90	100	110	120
PTVALYGLKQ	DWEGISSSSH	NDNHFNNKGY	SFKYENNPFL	GFAGAIGYSM	GGPRVEFEVS
130	140	150	160	170	180
YETFDVKNQG	NNYKNDAHRY	CALGOODNSG	IPKTSKYVLL	KSEGLLDISF	MLNACYDIIN
190	200	210	220	230	240
ESIPLSPYIC	AGVGTDLISM	FEATNPKISY	QGKLGLSYSI	NPEASVFIGG	HFHKVIGNEF
250	. 260	270	280	290	300
RDIPTLKAFV	TSSATPDLAI	VTLSVCHFGI	ELGGRENE		

Fig. 7B

10	20	30	40		
ATGAATTGCA	TTTTTAAAAA	TATABCAACT	₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩	50	60 CTTCTTACCT
70	BO	90	100		
GGAATATCAT		AGTACAGAAC		110	. 120
130	140	150	160		CTATATCAGT
GGGAAATATG	TACCAAGTGT			170	180
190	200	TTCACATTTT 210	220		
ACAACAACCG	GAGTATTTGG	ATTAAAGCAA	CATTCCCAMC	230	2.0
250	260	270	280		ATCTAAAAAT
TCTCCAGAAA	ATACATTTAA			290	300
310	320	330		AATATGAAAA	TAATCCATTT
CTAGGTTTTG			340	350	360
370	380	TGGTTATTTA			GTTAGAAATG
TCCTATGAAA			400	410	420
430	440	GAAAAACCAG		ATAAGAACGA	TGCTCACAAA
TATTATGCTT		450	460	470	480
490	500	CAGTGGGGGA	AAGCTAAGCA	ATGCAGGTGA	TAAGTTTGTT
TTTCTAAAA		510	520	530	540
550	560	ACTTGATATA	TCACTTATGT	TGAATGCATG	CTATGATGTA
ATAAGTGAAG		570	580	590	600
610	620	CTCTCCTTAC		GTGTTGGTAC	TGATTTAATA
TCCATGTTTG		630 CCCTAAAATT	640	650	660
670	680			GAAAGTTAGG	TTTGAGTTAC
TCCATAAGCC	CAGAAGCTTC	690	700	710	720
730	740			TTCATAAGGT	GATAGGGAAT
		750	760	770	780
790	BOO	TATGATACCC		CTCTCACAGG	TAATCACTTT
•		810	820	. 830	840
ACTATAGTAA 850	CACIMAGIGI	ATGCCACTTT		TTGGAGGAAG	GTTTAACTTT
TAA	860	870	880	890	900
	• • • • • • • • • • • • • • • • • • • •	••••••	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • •	••••

Fig. 8A

60	50	40	30	20	10
GVFSAKQERN	GKYVPSVSHF	DNVGGNFYIS	GISFSDAVQN	TLVSLMSFLP	MNCKKFFITT
120	. 110	100	90	80	70
MNGPRIELEM	LGFAGAVGYL	YSFKYENNPF	SPENTFNVPN	DWDGSTISKN	TTTGVFGLKQ
180	170	160	150	140	130
SLMLNACYDV	FLRNEGLLDI	KLSNAGDKEV	YYALTHNSGG	GNNYKNDAHK	SYETFDVKNQ
240	230	220	210	200	_. 190
GGHFHKVIGN	SISPEASVEV	SYQGKLGLSY	SMFEAINPKI	ICAGVGTDLI	ISEGIPFSPY
300	290	280	270	260	250
		GVELGGRENE	TIVTLSVCHF	STSTLTGNHF	EFRDIPAMIP

Fig. 8B

10	20	30	40	50	60
ATGGAAAATC		GAAAAACAAA	TTCTTTACAA	TAAGTACAGC	AATGGTATGC
. 70	80)	90	100	110	120
TTATTGTTAT	TACCTGGTAT	ATCATTTTCA	GAAACTATAA	ACAACAGTGC	TAAAAAACAG
130	140	150	160	170	180
CCTGGGTTAT	ATATCAGTGG	GCAGTACAAA	CCTAGTGTTT	CAGTTTTTAG	TAATTTTTCA
190	200	210	220	230	240
GTAAAAGAAA	CTAATGTTCC	CACAAAGCAG	TTAATAGCAC	TTAAAAAAAGA	CATTAATTCT
250	260	270	280	290	300
GTTGCAGTTG	GTAGTAATGC	TACTACAGGT	ATTAGCAATC	CAGGTAATTT	CACAATTCCT
310	320	330	340	350	360
TATACTGCAG	AATTTCAAGA	TAATGTTGCC	AATTTCAATG	GGGCTGTTGG	TTACTCTTTT
370	380	390	400	410	420
CCTGATAGTC	TAAGAATTGA	AATAGAGGGA	TTTCATGAAA	AATTTGATGT	CAAAAACCCT
430	440	450	460	47.0	480
GGAGGTTACA	CACAAGTAAA	AGATGCGTAC	CGTTATTTTG	CACTAGCACG	TGATTTAAAA
490	500	510	520	530	540
GATGGCTTCT	TTGAACCTAA	AGCGGAAGAT	ACAGGTGTTT	ATCATACTGT	TATGAAAAAT
550	560	570	580	590	600
GATGGATTAT	CTATTTTATC	TACTATGGTT	AACGTCTGTT	ACGATTTTC	TGTAGATGAA
610	620	630	640	650	660
TTACCAGTCT	TACCTTATAT	ATGTGCAGGT	ATGGGTATAA	ACGCCATAGA	ATTCTTCGAC
670	680	690	700	710	720
GCTTTACATG	TAAAATTTGC	TTACCAAGGC	AAACTAGGTA	TTAGCTATCA	
730					
AAAGTAAATT		TGGGTATTAC	CATCAAGTAA	TAGGCAATCA	
790					
				AAGTCACATC	
850	860	870	880	890	900
ACACTTGACA	TTGCATACTT	TGGTGGCGAA	GTTGGAATA	GATTCACATI	TTAA

Fig. 9A

	20	JU	40	50	60
MVCLLLLPGI	SFSETINNSA	KKQPGLYISG	QYKPSVSVFS	NFSVKETNVP	TKQLIALKKD
. 70	80	90	100	110	120
·INSVAVGSNA	TTGISNPGNF	TIPYTAEFQD	NVANFNGAVG	YSFPDSLRIE	IEGFHEKFDV
130	140	150	160	170	180
KNPGGYTQVK	DAYRYFALAR	DLKDGFFEPK	AEDTGVYHTV	MKNDGLSILS	TMVNVCYDFS
190	200	210	220	230	240
VDELPVLPYI	CAGMGINAIE	FFDALHVKFA	YQGKLGISYQ	LFTKVNLFLD	GYYHQVIGNQ
250	260	270	. 280	290	300
FKNLNVNHVY	TLKESPKVTS	AVATLDIAYF	GGEVGIRFTF		

Fig. 9B

10	20	30	40		
ATGATATATA	AAGAAAAACT	TACTAGAGTG	GGAGAATATA	50	60
70	б́в				TTTATCATTT
ATTCTTTTTT	CTTATATCTT	90	100	110	120
120		TCTAGTGCTG	GTAAATATTA	TTAGATATAA	CAGCCTTGCT
130	140	150	160	170	180
ATATGTGTTA	TCAGTCTACT	AAGAACTAAT	ATCTTTAACG	TTAGCACAAA	AAAATTAATA
190	200	210	220	230	
AAAGATAAAT	GTCGTGATAC	TAAGTTTAGT	AACATGAATT		240
250	260	270		GTTATTTGTA	CGGTAAACCG
TTAAATTTAC	AAATTTTTTA		280	290	300
310		TGGAATATTT	TCCTTTATTA	GAAACTTTCA	AAATAACACA
	320	330	340	350	360
CTAATAATTC	CTAATGATAG	TAAATGCGGC	TTCTATACCA	CGTTATGGGA	TAATCCAGCA
370	380	390	400		
CTACATTATA	CATATACACT	TACTGGCAGT		410	420
430	440		GAGTACCGTA	ATTTTTTTGA	CATTCTATAT
GAAAACATTA		450	460	470	480
		TAAATTACTT	ATTAACTATA	ACCGTTCTGT	ATTAAACCAA
490	500	510	520	530	540
CATAATAAAA	ATACTCTCGT	AATAATACCA	ATACCTAATG	CTAGAGAGTT	
550	560	570			CAGTAATGAA
ATTCGAGTAA	GGAATATATC		580	590	600
		**************************************	GAAAGTTCTT	ATGAGTGCTA	A

Fig. 10A

10	20	30	40	50	60
MIYKEKLTRV	GEYILAYLSF	ILSTYIFLVL	VNIIRYNSLA	ICVISLLRTN	IFNVSTKKLI
70.	80	90	100	110	120
KDKCRDTKFS	NMNCYLYGKP	LNLQIFYGIF	SFIRNFQNNT	LIIPNDSKCG	FYTTLWDNPA
130	140	150	160	170	180
LHYTYTLTGS	EYRNFFDILY	ENIICQCKLL	INYNRSVLNQ	HNKNTLVIIP	IPNAREFSNE
190	200	210	220	230	240
IRVRNISINK	ESSYEC				

Fig. 10B

10	20	30	40	50	60
ATGAATAAAA 1	AAAACAAGTŢ	TATTATAGCT	ACAGCATTGG	יים איים מיים מיים מיים מיים מיים מיים מ	60
, 0	80	90	100	110	100
AGTGTATCGT	TTTCAGAGGT	TACAAACAGC	AGTATTAAAA	AACACTCTCC	120
130	140	150	160	170	100
AGTGGACAAT A	ACAAACCAAG	TGTTTCTGTT	TTTAGTAGTT	ע ע שייי ע אייי איי	180
190	200	210	220	220	0.40
ACTATCACAA A	AAAATCTTAT	AGCGTTAAAA	AAAGATATTA	בסט	240
250	260	270	280	200	200
GATGCTAGTC A	AAGGTATTAG	TCATCCAGGA	AATTTTACTA	TACCTTATAT	300
210	320	330	340	250	3.50
GAAGATAATG (CTTTTAATTT	CAACGGTGCT	ATTGGTTACA	ТТАСТСААСС	UOC.
370	380	390	400	410	400
GAAATAGAAG	STTCCTATGA	AGAATTTGAT	GCTGAAAACC	רייההאההיייא	42U
430	440	450	460	470	
GATGCCTTTC G	GGTACTTTGC	TTTAGCACGT	GATATGGAAA	OUT E	40U
490	500	510	520	F 2 0	540
GCACAAAGCT C	CAC			5,50	540

Fig. 11A

10	20	30	40	50	60
MNKKNKFIIA	TALVYLLSLP	SVSFSEVTNS	SIKKHSGLYI	SGQYKPSVSV	FSSFSIKETN
70	. 80	90	100	110	120
TITKNLIALK	KDINSLEVNA	DASQGISHPG	NFTIPYLAAF	EDNAFNFNGA	IGYITEGLRI
130	140	150	160	170	180
EIEGSYEEFD	AENPGGYGLN	DAFRYFALAR	DMESNKFLPK	AOSS	

Fig. 11B

10	20	30	40	50	60
TCTAGAATAC	ATGATGAAAA	TTATGCTATT	ACAACAAATA	ATAAATTATC	CATCGCATCT
70	80	90	100	110	. 120
ATTATGGTTA	ACACCTGCTA	TGATATTTCA	ATTAATAATA	CATCAATAGT	ACCGTATTTA
130	140	150	160	170	180
TGCACAGGCA	TTGGTGAAGA	TCTTGTAGGG	CTTTTTAATA	CAATACATTT	TAAACTTGCA
190	200	210	220	230	240
TATCAAGGGA	AAGTTGGAAT	GAGTTATTTG	ATAAATAACA	ATATCCTATT	ATTTTCTGAC
250	260	270	280	290	. 300
ATATATTATC	ATAAAGTCAT	GGGTAACAGA	TTTAAAAATT	TGTACATGCA	ATATGTAGCT
310	320	330	340	350	360
GATCCTAATA	TTTCTGAAGA	AACTATACCT	ATATTAGCAA	AACTTGATAT	TGGTTATTTT
370		390	400		420
GGAAGTGAAA	TTGGAATAAG	GTTTATGTTT	AACTAA	• • • • • • • •	

Fig. 12A

60	50	40	30	20	10
LENTIHEKLA	CTGIGEDLVG	INNTSIVPYL	.IMVNTCYDIS	TTNNKLSIAS	SRIHDENYAI
. 120	. 110	100	. 90	80	70
ILAKLDIGYF	DPNISEETIP	FKNLYMQYVA	IYYHKVMGNR	INNNILLFSD	YQGKVGMSYL
180	170	160	150	140	130
				N	GSETGTREME

Fig. 12B

				_	
10	20	30	40	50	60
ATGACAAAGA	AATTTAATTT	TGTAAATGTT	ATATTAACAT	TTTTGTTATT	TCTTTTCCCA
70	80	` 90	100	110	120
CTTAAGTCAT	TTACAACATA	TGCAAATAAT	AACACAATCA	CTCAAAAAGT	TGGATTGTAC
130	140	150	160	170	180
ATAAGTGGTC	AATATAAGCC	AAGTATTCCT	CATTTCAAGA	ATTTTTCAGT	AGAAGAAAAT
190	200	210	220	230	240
GACAAAGTAG	TAGATTTGAT	AGGTCTTACA		CATATATCAC	AGAACATATA
250	260	270	280	290	300
TTACGAGATA	ATACAAAATT	CAACACTCAT	TATATTGCAA	AGTTCAAGAA	CAATTITATA
310	320	330	. 340	350	360
AATTTCAGCA	GTGCAATTGG	TTATTATTCT	GGGCAAGGAC	CAAGGTTAGA	AATAGAAAGC
370	380	390	400	410	420
TCTTATGGGG	ATTTTGATGT	TGTAAATTAT	AAAAATTATG	CAGTACAAGA	TGTTAATAGA
430	440				
TATTTTGCTT	TAGTACGTGA	AAAAAATGGT	TCAAATTTCT	CTCCAAAACC	ACATGAAACT
490	500	510	520	530	540
AGTCAACCCT	CTGACAGTAA	TCCTAAAAA	TCTTTTTATA	A CTTTAATGAA	GAATAATGGG
550	560	570	580	590	600
GTATTTGTTG	CATCAGTAAT	AATCAACGGT	TGTTATGAT	r TTTCTTTTAA	TAACACAACA
610	620				
ATATCACCTI	ACGTATGTA	R AGGAGTTGG	A GGAGATTTT	A TAGAGTTTT	TGAAGTAATG
670	680			-	
CATATCAAGT	TTGCTTGCC	A AAGTAAGGT	T GGTATTAGC	T ATCCAATAT	TCCCTCTATT
730	74				
ACTATTTTT	CTGATGCAC	A TTATCACAA	G GTCATAAAT	A ATAAATTA	A CAACCTACAT
790	08 0				=
GTTAAGTAT	r CATATGAAC	T TAAAAACTC			C AGCCAAACTA
85	0 86	0 87	0 88	10 89	0 900
AACATTGAA	T ATTTTGGTG	G TGAAGTTGG	G ATGAGATTI	A TATTTTAA.	

Fig. 13A

60	5.0	40	30	20	10
HFKNFSVEEN	ISGQYKPSIP	NTITOKVGLY	LKSFTTYANN	ILTFLLFLFP	
120	110	100	90	80	70
GQGPRLEIES	NESSAIGYYS	YIAKFKNNFI	LRDNTKFNTH	TDVTYITEHI	DKVVDLIGLT
180	170	160	150	140	130
SFYTLMKNNG	SQPSDSNPKK	SNFSPKPHET	YFALVREKNG	KNYAVQDVNR	SYGDEDVVNY
240	. 230	220	210	200	190
GISYPISPSI	HIKFACQSKV	GDFIEFFEVM	ISPYVCIGVG	CYDESENNTT	VFVASVIING
300	290	280	270	. 260	250
MRFIF	NIEYFGGEVG	PTITSATAKL	VKYSYELKNS	UTNNKENNLH	TEN DAUVUK

Fig. 13B

					60
. 10	20	30	40	50	
ATGAGCAAAA	AAAAGTTTAT	TACAATAGGA	ACAGTACTTG	CATCTCTATT	120
70	ลก	` 90	. 100	110	
TCTATTGAAT	CCTTTTCAGC	TATAAATCAT	AATCATACAG	GAAATAACAC	180
130	140	150	160	170	. 100
TATATTACAG	GGCAGTATAG	ACCAGGAGTA	TCCCATTTA	GCAATTTCTC	240
100	200	210	220	230	2.0
ACTAATGTTG	ATACAATACA	ACTAGTAGGA	TATAAAAAAA	GTGCGTCTTC	300
250	260	270	280	290	500
AACACTTATT	CAAACTTTCA	AGGTCCATAT	ACTGTTACAT	TTCAAGATAA	TGCTGCTAG1
210	320	330	340	350	300
TTCAGTGGAG	CAATTGGATA	TTCTTACCCC	GAAAGTCTAA	GACTTGAACT	TGAAGGTTCT
270	380	390	400	1 410	420
TACGAAAAAT	TTGATGTCAA	AGATCCTAAA	GACTACTCA	CAAAAGATGC	TTTTAGGTTT
430	440	450	460) 4/0	400
TTTGCTCTAG	CACGTAATAC	GTCTACTACT	GTTCCTGAT	G CTCAAAAATA	TACAGTTATG
400	5.00	510	52) 530) 340.
AAGAATAAT	GCTTATCTG	TGCATCAATC	ATGATCAAT	G GTTGTTATG!	TCTATCTTTT
CE	56	n 570) .58	0 591) 050
AATAATTTA	TCGTATCAC	C TTATATATG	r gcaggtatt	G GTGAAGATT	CATTGAATTT
	. 62	ი 630	ი 64	0 65	0 000
TTTGATACT	T TGCACATTA	A ACTTGCTTA	T CAAGGAAAA	C TAGGTATTA	G TTATTACTTC
67	a 69	ი 69	ი 70	10 11	0 720
TTTCCTAAG	A TTAATGTAT	T TGCTGGTGG	G TACTATCAT	A GAGTTATAG	G GAATAAATTT
77	n 74	เก 75	0 /1	າ ບ	0
דדדממממ	A ATGTTAACO	A TGTTGTTAC	A CTTGATGA	AT TTCCTAAAG	C AACTTCTGCA
~	.c 81	۱n 81	.ი 8:	20 83	30 040
CTACCTAC!	C TTAATGTT	C TTATTTTGG	T GGTGAAGC	IG GAGTAAAG	TACATTTAA
8!	_		70 8	80 89	900
					• • • • • • • • • • • • • • • • • • • •

Fig. 14A

60	50	40	30	20	10
SHFSNFSVKE	YITGQYRPGV	NHTGNNTSGI	SIESFSAINH	TVLASLLSFL	MSKKKFITIG
	110	100	90	80	70
ESLRLELEGS	FSGAIGYSYP	TVTFQDNAAS	NTYSNEQGPY	YKKSASSIDP	TNVDTTOLVG
180	170	160	150	140	130
MINGCYDLSF	KNNGLSVASI	VPDAQKYTVM	FALARNTSTT	DYSAKDAFRE	YEKEDVKDPK
	230	220	210	200	190
YYHRVIGNKF	FPKINVFAGG	OGKLGISYYF	FDTLHIKLAY		
	290			260	250
,		GEAGVKETE.	VATLNVAYFG		

Fig. 14B

				5.0	60			
10	20	30	40	50				
ATGF.GTGCTA	AAAAAAAGCT			TAGTATGTTT	AGTGTCATAC 120			
70	80	90	. 100	110				
TTACCTACTA	AATCTTTGTC			ATAACACTAA	180			
130	140	150	160	170				
	GTGGACAATA				240			
190	200	210	220	230				
GAAACTTATA	CTGACACTAA			AAGATATTAA	300			
250	260	270	280	290				
GATATAACAA	САААТААААА			CAAAATTTCA	AGATAATGCT			
310	320	330	. 340	350	360			
GTTAGCTTCA	GTGCAGCTGT			GTCCAAGGGT	TGAGGTAGAA			
370	380	390	400	410	420			
TGGTCTTATG	AAGAATTTGA	CGTTAAAAAT		ACGTAGTAAG	TGAAGCCTTC			
430	440	450	460	47.0	480			
AGGTATATTG	CTTTAGCAAG	AGGAATTGAT			AACAAATAAG			
490	500	510	520	530				
TATGTTGTTA	TAAAGAACAA	TGGCTTATCT	GTCGCATCCA		TGGCTGTTAT			
550			580		•			
GATTTTTCTI	TAAACAATTT	AAAAGTATCA			TGGTGGGGAC			
610								
ATTATAGAAT	TTTTTAGTGC	TGTAAGTTTT			GGTAGGTATC			
670					=			
AGTTATCCAT	TATTCTCTA	TATGATTATA			A TAAGGTCATA			
. 730					-			
GGAAATAAA!	TTAACAATT	AAATGTTCA	CACGTTGTT	A GTCTTAACA	G TCATCCTAAG			
79								
TCTACTTTT	G CAGTAGCTA			G GTAGTGAAT	T TGGGTTAAAA			
85	0 86	870	88	0 89	0 900			
TTTATATTT	T AA		·					
Fig. 15A								
10	20	30	40	50	60			
IV	CONTROLIS	T.PTKSI.SNLN	NTNNNTKCTG	LYVSGOYKPT	VSHFSNFSLK.			
		90	100	110	120			
70	00 תרסעדתעג זה	DITTINKKENI			SQDSPRVEVE			
ETITOTKELL	140	150	160	170	180			
. T20	DCMVINICENE 140	בבט חדמת מדעק			VASILINGCY			
	JUU THACANTARA	210	220	230	240			
190	LUV LUVICECCE	TTEFFERIUS	KEAYOGKVG		FADGYYHKVI			
		270	28	290	300			
250		. 210 2117 Carry 12			• ••••••			
GNKFNNLNV(1 HAAPTNRHEL	/ SIEWAWITH/	Eligation					

Fig. 15B

10	20	30	40	50	60
ATGAGTAAAA	TATTTTAAAA	TACAATAGGA	GCAACACTTA	TTCATATGTT	GTTACCTAAC
70	80	. 90	100	110	. 120
ATATCTTTTC	CAGAAACTAT	TAACAATAAC	ACTGATAAAC	TTTCTGGGTT	ATATATAAGT
130	140	150	160	170	180
GGGCAATATA	AACCAGGGAT	TTCTCATTTC	AGCAAATTTT	CAGTCAAAGA	AATCTATAAT
190	200	210	220	230	240 -
GATAACATTC	AACTAATTGG	GTTAAGACAC	AACGCAATTT	CTACTAGTAC	CCTTAATATT
250	260	270	280	290	300
AATACAGATT	TTAATATCCC	CTATAAAGTA	ACATTTCAAA	ATAACATTAC	CAGCTTTAGT
310	320	330	340	. 350	360
GGAGCTATTG	GTTATTCTGA	TCCCACAGGG	GCAAGATTTG	AGCTTGAAGG	TTCTTATGAA
370	380	390	400	410	420
GAATTTGATG	TGACAGATCC	TGGAGACTGC	TTAATAAAAG	ATACCTATAG	ATATTTCGCT
430	440	450	460	470	480
TTAGCTAGAA	ACCCATCAGG	TTCTAGCCCT	ACCTCAAACA	ACTATACTGT	TATGAGAAAT
490	500	510	520	530	540
GATGGTGTTT	CCATTACTTC	TGTTATATTI	AATGGCTGTT	ATGACATCTI	TTTAAAGGAT
550	560	570	580	590	600
TTAGAAGTAT	CACCTTATG	ATGTGTTGG	GTAGGTGGA	ATTTTATAGA	ATTTTTTGAC
610	620	630	640	650) 660
GCATTACAC	TTAAATTAG	ATACCAAGG	C AAGTTAGGT	A TCAATTATCI	A CTTATCGACT
670	680	69	0 70	0 710	720
CAAGCAAGC	G TATTTATTG	A TGGATATTA	T CATAAGGTT	A TAGGAAATC	A ATTCAACAAT
73	0 74		-	-	=
CTAAATGTT	C AACACGTGG	C TAGTACAGA	T TTTGGACCT	G TATACGCAG	T AGCCACACTT
79					0 840
AACATTGGT	T ATTTTGGTG	G TGAAATCGG	A ATTAGACTI	A CATTTTAA.	
			Fig. 16A		
1	.0 2			. •	60
MSKKNFITI	G ATLIHMLLE	N ISFPETIN	N TOKLSGLY	IS GQYKPGISH	F SKFSVKEIYN
-	in F	20	90 10	00 13	120
DNIOLIGLE	H NAISTSTL	I NTDFNIPY	KV TEQNNITS	FS GAIGYSDPT	G ARFELEGSYE
17	RO 14	10 1:	50 1	PO 1	70 200
EFDVTDPGI	C LIKDTYRY	A LARNPSGS	SP TSNNYTVM	RN DGVSITSV	IF NGCYDIFLKD
10	an '21	00 . 2	10 2	20 23	30 240
LEVSPYVC	VG VGGDFIEF	FD ALHIKLAY	QG KLGINYHL	ST QASVFIDG	YY HKVIGNQFNN
. 2	50 . 2	60 2	70 2	.80 2	90 300
LNVQHVAS	TD FGPVYAVA	TL NIGYFGGE	IG IRLTF		••.••••
_					

Fig. 16B

. 10	20	30			
ATGAATAATA	GAAAAAGTTTT	טט. במבנות לוחיתייתיים	40	50	60
70	8.0	TITTATAATA	GGTGCATCAT	TACTAGCAAG	60 CTTATTATTC
130	140	AGGAAATGTA	AGTAACCATA	CTTATTTTAA	ACCTAGGTTA
	~30	1 711	1/0		
190	200	ACCAGGAGTT	TCTCATTTTA	GCAAATTTTC	180 AGTCAAAGAA
250	260	ACIAGITGGG	CTTAAAAAGG	ACATCAGTGT	CATAGGGAAC
310	320	AAATTTCAAC	TTTCCTTACA	TTGCAGAATT	TCAAGACAAT
370	380	AATTGGATAC	TTGTATTCCG	AGAATTTTAG	AATTGAAGTA
430	440	TGATGTTAAA	AATCCAGAAG	GATCTGCTAC	AGACGCATAC
490	500	IGCTATGGAT	GGCACTAATA	AATCTAGTCC	TGATGACACA
550	560	AAATGACGGG	TTATCAATTT	CATCAGTAAT	GATAAATGGG
610	620	IGATATACCA	GTAGTACCGT	ATGTATGCGC	AGGAATAGGA
GGAGATTTCA 670	680	TAATGATTTA	CATGTTAAGT	TTGCTCATCA	AGGCAAGGTA
GGTATTAGTT	סטט איירייי אייאייר א	690	700	710	720
GGTATTAGTT 730	740	CCCTGAAGTA	AGTTTATTTC	TTAACGGATA	TTACCATAAA
GTAACAGGTA	ACACATTA A	750	760	770	780
GTAACAGGTA 790	800	HAACTTACAC	GTTCAACACG	TAAGTGATTT	AAGTGACGCT
CCTAAGTTCA 850	860	TGCTACACTC	AATGTTGGGT	ACTTTGGTGG	CGAAATTGGA
GTAAGATTTA		870	880	890	900
		••••••	• • • • • • • • •	• • • • • • • • • • • • • • • • • • • •	

Fig. 17A

			40 SNHTYFKPRL 100		
EASYEEFDVK 190	NPEGSATDAY	RYFALARAMD	FPYLAEFQDN 160 GTNKSSPDDT	AISFSGAIGY 170 RKFTVMRNDG	LYSENFRIEV 180 LSISSVMING
CYNFTLDDIP 250 VTGNRFKNLH	VVPYVCAGIG 260	GDFIEFFNDL			240 SLFLNGYYHK 300

Fig. 17B

10	20	30	40	50	60
TAGCAGCACT	AAAAAACAGT	TTGGGTTATA	TGTTAGTGGA	CAACACCAGC	CTAGTGTTTC
70	68	90	100	110	120
TATTTTTAGC	AATTTCTCAG	TAAAGGAAAC	TAATTTTCCT	ACAAAGTATT	CTAGCAGCTT
130	140	150	160	170	180
CTTAAAAAAA	GACATTAATT	CTGTCGAATT	TGACGATAGT	GTTACTGCTG	GCATTAGTTA
190	200	210	220	230	240
CCCACTTAAT	TTCAGTACTC	CTTATATAGC	TGTATTTCAA	GATAATATTT	CTAATTTTAA
250	260	270	280	290	300
TGGCGCTATT	GGGTACACTT	TTGTTGAAGG	CCCAAGAATT	GAAATAGAAG	GTTCTTATGA
310	320	330	. 340	350	3,60
AGAATTCGAT	GTCAAAGACC	CTGGAAGATA	TACAGAAATA	CAAGATGCAT	ACCGTTACTT
370	380	390	400	410	420
TGCTTTAGCA	CGTGATATAG	ACTCTATTCC	TACTAGCCCA	AAAAATAGAA	CTTCACATGA
430	440	450	460	470	480
TGGCAACAGT	TCATATAAGG	TATACCACAC	TGTAATGAAA	AATGAAGGAC	TATCTATAAT
490	500	510	520	530	540
ATCCATTATG	GTCAATGGCT	GCTATGATTT	TTCTTCAGAT	AATTTATCAA	TATTACCTTA
550	560	570	580	590	600
TGTATGTGGT	GGTATAGGTG	TAAATGCTAT	AGAGTTTTTC	GATGCATTAC	ATGTTAAATT
610	620	630	640	650	660
CGCGTGTCAG	GGTAAATTAG	GTATTACTTA	TCCATTATCT	TCCAACGTTA	GTTTATTTGC
670	680	690	700	710	720
TGGTGGATAT		TAATGGGCAA	CCAATTTAAA	AATCTAAATG	TTCAACATGT
730	740	750	760		780
AGCTGAACTT				GCTACACTTG	ACATTGGGTA
790		810		. 830	840
TTTTGGTGGT	GAAATTGGAG	CAAGGCTTAT	ATTTTAA	•••••••	

Fig. 18A

60	50	40	. 30	.20	10
DDSVTAGISY	LKKDINSVEF	NEPTKYSSSE	IFSNFSVKET	VSGQHQPSVS	SSTKKQFGLY
120	110	100	90	80	70
TEIQDAYRYF	EFDVKDPGRY	PRIEIEGSYE	GAIGYTFVEG	VEQDNISNEN	PLNFSTPYIA
180	170	160	150	140	130
SSDNLSILPY	SIMVNGCYDF	VMKNEGLSII	GNSSYKVYHT	TSPKNRTSHD	ALARDIDSIP
240	230	220	210	200	190
OEKNINAOHA	GGYYHQVMGN	PLSSNVSLFA	ACQGKLGITY	EFFDALHVKF	VCGGIGVNAI
300	290	280	270	260	250
		F	FGGEIGARLI	SAVATLDIGY	AELNDAPKVT

Fig. 18B

10	20	30	40	50	60
	AAAGATTTT				TTTCTTACCT
70	80.	90	100	110	120
AGCGTATCTT	TTTCTGAATC	AATACATGAA	GATAATATAA	ATGGTAACTT	TTACATTAGT
130	140	150	160	170	180
GCAAAGTATA	TGCCAAGTGC	CTCACACTTT	GGCGTATTTT	CAGTTAAAGA	AGAGAAAAAC
190	200	210	220	230	240
ACAACAACTG	GAGTTTTCGG	ATTAAAACAA	GATTGGGACG	GAGCAACAAT	AAAGGATGCA
250	260	270	280	290	300
AGCAGCAGCC	ACACAATAGA	CCCAAGTACA	ATATTCTCCA	TTTCAAATTA	TTCATTTAAA
310	320	330	340	350	360
TATGAAAACA	ATCCATTTTT	AGGGTTTGCA	GGAGCTATTG	GCTACTCAAT	GGGTGGTCCA
370	380	390	400	410	420
AGGGTAGAGT	TTGAAGTGTC	TTACGAAATA	TTTGATGTAA	AAAACCAAGG	TAACAGTTAC
430	440	450	460	470	480
AAGAACGATG	CTCACAAATA	TTGCGCTTTA	TCAAGACACA	CCGGAGGTAT	GCCACAAGCC
490	500	510	520	530	540
GGTCATCAAA	ATAAATTTGT	CTTCCTAAAA	AATGAAGGAT	TACTTGACAT	ATCACTTATG
550	560	570	580	590	600
ATAAACGCAT	GTTATGATAT	AACAATCGAC	AGCATGCCAT	TTTCTCCATA	TATATGTGCA
610	620	630	640	650	660
GGTATTGGTA	GTGACTTAGT	TTCGATGTTT	GAAACTACAA	ATCCTAAAAT	
670	680	690			•
GGAAAATTAG	GTGTAAGTTA	CTCCATAAGC		CTGTTTTTGI	
730		750			
TTTCACAGAG				CAATAACTC	
790					
ACAGAAATTA					A CTTCGGACTA
850		•	886	890	900
GAGCTTGGAG	GCAGGTTTAC	TTTTTAA	• • • • • • • • • • • • • • • • • • • •		

Fig. 19A

60	50	40	30	20	ΤO
GVFSVKEEKN	AKYMPSASHF	DNINGNEYIS	SVSFSESIHE	ALISLMSFLP	MNCKREFIAS
120	110	100	90	80	70
GAIGYSMGGP	YENNPFLGFA	IFSISNYSFK	SSSHTIDPST	DWDGATIKDA	TTTGVFGLKQ
180	170	. 160	150	140	130
NEGLLDISLM	GHQNKFVFLK	SRHTGGMPQA	KNDAHKYCAL	FDVKNQGNSY	RVEFEVSYEI
240	230	220	210	200	190
PEASVEVGGH	GKLGVSYSIS	ETTNPKISYQ	GIGSDLVSMF	SMPFSPYICA	INACYDITID
300	290.	280	270	260	250
	ELGGRETE.	VITATCHECT.	TEIKGTOFTT	DIPATTPAGA	FHRVIGNEFK

Fig. 19B

	60	50	40	· 30	20	10
	CTTTACACAT	TATTAACTTC	GCATTAGTAT	TACAGTAACT	AAAAAACTTT.	ATGAAATATA
	120	110	100	90 -	80	70
	CATTAGTGGA	ACAACTTCTA	AGTACAATTC	AGCACGTGCC	TTTATAGTCC	TTTATACCTT
	180	170	160	150	140	130
	ACAAAGTTTT	CTAAAGAAGA	ATTTTTTCAG	ACATTTTGGA	CAACAGCGTC	AAATATATGC
!	240	230	220	210	200	190
٠.	CAATAATGAT	ATATTATAA	TTATCACATA	AGATCAACGA	TAGTTGGGTT	ACTAAGGTAT
١.	300	290	280	270	260	250
k.	CCCATTTCTA	ACAAAAATAA	TCATTTAAAT	TCAAAATTAT	GTCTTAAGGT	ACAGCAAAGA
į	360	350	. 340	330	320	310
L	AGAAGTATC	GAATAGAACT	GGCAATTCAA	TTATTCAATA	GAGCTATTGG	GGATTTGCAA
)	420	410	400	390	380	370
:	TCACAAATA	TAAATGACTC	AACAATTATT	AAACCCAGGA	TTGATACTAA	CATGAAATAT
)	`48(470	460	. 450	440	430
C	TTGGTACAC:	ATAGCGGAGA	AGTGATGGAA	TCACATATGC	CTCATGGAAG	TGCGCTTTAT
)	.54	530	520	510	500	490
3	CTCATTTAT	TACTTGACGT	AATGAAGGTT	ACTTCTGAAA	ATAAGTTTGT	GCAAAAACTG
)	60	590	580	570	560	550
A	TATATGTGC	TTTCACCTTA	AAAATGCCTT	AACAACTGAA	GTTATGACAT	TTAAACGCAT
	66	650	€40	630	620	610
		AAAACAAAAT	GAGACAACAC	ATCTATGTTT	CTGATCTCAT	GGTATTGGTA
_		710	. 700	690	680	670
		CTGTTTTTGC	TCAAGAGTTI	TACTATAAAC	GTTTAAACTA	GGAAAGTTAG
_			760	750		730
		CTCTATTACC			TAATAGGTAA	TTTCATAAAG
_				810		790
		TGTGCCATTI			TACAACAGTO	AACATTAAAG
O	90			870	860	850
•				TTAA	GATTTTCTT	ATTGGAAGTA

Fig. 20A

∴60	50	40	30	. 20.	10
IFSAKEEOSF	KYMPTASHFG			ALVLLTSFTH	MKYKKTETVT
120	110	100	90	80	70
GNSRIELEVS	GFARAIGYSI	SEKYKNNPEL		LSHNIINNND	. •
180	170	160	150	140	130
NEGLLDVSFM	AKTOKEVLLK	SDGNSGDWYT	CALSHGSHIC	NNYLNDSHKY	HEIFDTKNPG
	230	220	210	200	190
SRVSVFAGGH	GKLGLNYTIN	ETTQNKISYQ	GIGTDLISME	KMPESPYICA	LNACYDITTE
300				260	250
•••••	igsrfff	TLDVCHFGLE	NIKVQQSATV	GIPTLLPDGS	FHKVIGNEFK

Fig. 20B

	20	30	40	. 50	60
10	. 20 בתמתמתמתמת			TTGCACTTCC	ACTATTGTTA
70	80	90	100	110	120
	ACTATTTAG	GTGTAATATG	AATTGCAAAA	AAATTCTTAT	AACAACTGCA
130	140	150	160	170	. 180
TTAATATCAT	TAATGTACTC	TATTCCAAGC	ATATCTTTTT	CTGATACTAT	
190	200	210	220	. 230	240
AACATGGGTG	GTAACTTCTA			CAAGTGTCTC	ACATTTTGGT 300
250	260	270		290	
				TTTTTGGATT	360
310	320	. 330	340		
		TAAGAATAAA 390	CACGCTGACT	TTACTGTTCC 410	420
3.70	380			CTATCGGTTA	CTCAATGGGT
TTCAGATACG	440	450	•		480
				ACGTAAAAAG	TCCTAATATC
490	500	510			540
AATTATCAAA	ATGACGCGCA	CAGGTACTGC	GCTCTATCTC	ATCACACATC	GGCAGCCATG
550	560	570	580	590	600
GAAGCTGATA	AATTTGTCTT	CTTAAAAAAA	GAAGGGTTA	TTGACATATC	
610	620	630			
AATGCATGTT				CTCCTTATAT	
670			-	•	
				C CTAAAATTTC	
730	740			G TTTTCATCG	
AAACTGGGCF				0 830	
					TAACTCAACT
850	860	87	0 88	0 899	ე 900
ACAATAAGT	GACCACAAT	TGCAACAGT	A ACACTAAAT	G TGTGTCACT	r TGGTTTAGAA
910	920	93	0 94	0 95	0 960
CTTGGAGGA	A GATTTAACT	r CTAA		• • • • • • • • • • • • • • • • • • • •	
	_				
]	Fig. 21A			
		_	_	. 54	5 60
10) 20	3	0 4	_	
MEYTNIYIL	CIYFALPLL	LIYEHYERCM	M NCKKILITI	N LISIMISIE	S ISFSDTIQDG
70) 8(9 9	e michect.KE	n wnespiikn	K HADETVPNYS
NMGGNFYIS	AIVESVORES	s seanneesn N 15	io 14GVEGIRA	50 17	0 180
TOVENINDET.	E FAGATGYSM	G GPRIEFEIS	Y EAFDVKSPI	II NYQNDAHRY	C ALSHHTSAAM
1.0	n. 20	0. 21	10 23	20 .23	0 240
EDUKEUFI K	N EGLIDISLA	I NACYDIINI	OK VPVSPYIC	AG IGTDLISME	E ATSPRISIQG
25	ი 26	n · 2°	70 2	80 25	0. 300
KLGISYSIN	P ETSVFIGGH	F HRIIGNEF	RD IPAIVPSN	ST TISGPOFAT	TLNVCHFGLE
31	0 32	.0 3:	30 3	40 35	360
LGGRFNF					

Fig. 21B

					cò
10	20	30	40	50	60
ATGAATTGCA	AAAAAATTCT '		GCATTAATGT	CATTAATGTA	CTATGCTCCA
70	80	90	100	110	120
AGCATATCTT	TTTCTGATAC		GATAACACTG		180
130	140	150	160	170	
AAATATGTAC					240
190	200	210	220	230	
ACTGTTGGAG			TGGAATGGAG		300
250	260	270	280	290	
CCAGAAAATA			TCGTTTAAAT	ACGAAAACAA	360
310	. 320	330	. 340	350	• • • • • • • • • • • • • • • • • • • •
GGGTTTGCAG			GGTGGCCCAA	0.2	TGAAGTTCTG 420
370	380	390	400	410	
TACGAGACAT					ACACAGATAC 480
430	440	450	460	470	
				CCGCAAGTAA	CAAATTTGTT 540
490	500	510	520	530	• • •
					CTATGACATA 600
550		570	580		
ATAATTGAAG					TGATGTTGTT 660
610					
					ATTAGGTTAT 720
670				, , , , ,	,
			GGTGGACACI		
730					,
					A AAACCAATTT
790				•	S ATTTAACTTC
850		870	, 00		
TGA					
		Fiş	g. 22A		
10	20	30	4.0	50	60
					VFSAKEERNS
70				110	120
					GGPRIELEVL
130	•				
					SFMINACYDI
190					
					I GGHFHRVIGN
	260		28		
		•	F GLELGGREN		

Fig. 22B

• •	20	30	40	50	60
. 10	20		GCATTGATAT		• •
•	AAAAAGIIII		100	110	120
70	• •		AACAGTATGT		TTACATATCA
130	140	150	160	170	180
			GGAATTTTTT		
190	200	210	220	230	240
		+	AAACTGGCAG		ATCTAGTCAA
250	260	270	280	290	300
			TACTCATTCA		CAACAAGTTT
AGICCAGAIG	320	330	340	350	360
			ATAGGCAGTC	•	AGTTGAGATG
370	380	390	400	410	420
TCTTATGAAG	•		GGTGATAATT	ACAAAAACGG	TGCTTACAGG
430	440	450	460	470	480
TATTGTGCTT				TGACTAGTGC	AACTGACAAA
490	500	510		530	540
TTTGTATATT	• • •		AACATATCAT	TTATGACAAA	CATATGTTAT
550					•
			CCTTACATAT		TGGTACTGAT
610					
		,	AAAATTTCTT	ATCAAGGAAA	GCTAGGGTTG
670					
GCCTACTTC		A GTCTTCGGTT	TCTTTTGGT	A TATATTTC!	A TAAAATTATA
730					
AATAATAAG	TTAAAAATG	TCCAGCCAT	G GTACCTATT	A ACTCAGACG	A GATAGTAGGA
790					
CCACAGTTT	CAACAGTAA	ATTAAATGT	A TGCTACTTT	G GATTAGAAC	T TGGATGTAGG
850	_				
TTCAACTTC	AA				

Fig. 23A

TO	20	30	40	50	60
MNCKKVFTIS	ALISSIYFLP	NVSYSNPVYG	NSMYGNEYIS	GKYMPSVPHF	GIFSAEEEKK
70	80	90	100	110.	120
KTTVVYGLKG	KLAGDAISSQ	SPDDNFTIRN	YSFKYASNKF	LGFAVAIGYS	IGSPRIEVEM
130	140	150	160	170	180
SYEAFDVKNP	GDNYKNGAYR	YCALSHODDA	DDDMTSATDK	FVYLINEGLL	NISFMTNICY
190	200	210	220	230	240
ETASKNIPLS	PYICAGIGTD	LIHMFETTHP	KISYQGKLGL	AYFVSAESSV	SFGIYFHKII
250	260	270		290	300
NNKFKNVPAM	VPINSDEIVG	POFATVTLNV	CYFGLELGCR	FNF	• • • • • • • •

Fig. 23B

	20	30	40	50	60
. 10	20	#2##2#################################	アクサヤスシャインス	CACTAACAAT	TCTTTTACCT
	AAAAATTTCT 80	TATAACAACI	100	110	. 120
· 70	222222	TAMBONTON	льсватаста	CAGGAAACTT	TTACATTATT
	140	150	160	1/0	
130	140	TOUR CACHE	CCCAACTTT	CAGCTAAAGA	AGAAAAAAAC
		210	220	230	240
190	200	712 442444444		GTGGTATCAT	CCTTGATAAA
	GAATTTTTGG 260		280	290	300
250	UOS Mara ammen	CCC		ATGAAAATAA	TCCATTTTTA
		330	340	350	360
310	32U	סכב בייבביית הייים י		GAATAGAATT	TGAAGTATCA
			400	410	420
370	300	, , ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		ACAATGATGC	ACATAAGTAT
			460	470	480
430	440	, ~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~			TTTTCTCAAA
			52	530	540
490	500 7	יבט די איזייטאריירטיינע			AATAAACAAA
			58	0 590	600
55(7 mmmcrccmm	0 7 CATATCTGC	A GGCATTGGT	A CTGACTTAA	ATTCATGTTT
			n 64	0 65	660
61	02 2	כ יינכריייאיירא מריייאיירא			A TCCAATAAGC
~ 7	^ 68	n 69	0 70	10 17	,,,,
7.0	00 	T GEGTGTGCA	C TTTCACAA	G TAACAAACA	A CGAGTTTAGA
	a 74	ın 75	a 76	วับ //	0 .55
13	יי אמייים אמריינים	C TEGAGGACT	C GCTCCAGA	TA ATCTATTTG	C AATAGTAAAG
70	n Rí	าก 81	.0 87	20 83	0.0
/5	ים כתכאייייייני	C GTTAGAAT!	TT GGGTACAG	GG TCAGTTTT	'A A
TTGAGTATA	T GICALLIA	30 0111.01			
			Fig. 24A		
					غ
1	LO :			30	60
MNCKKFLI	TT TLVSLTIL	LP GISFSKPI	HE NNTTGNFY	II GKYVPSISI	F GNFSAKEEKN
•	70	80	90 1	.00 • 13	10 . 120
TTTGIFGL	KE SWTGGIIL	DK EHAAFNIP	NY SFKYENNE	FL GFAGVIGY	SI GSPRIEFEVS
13	30 1	40 . 1	50 1	.60 1	70 180
YETFDVQN	PG DKFNNDAH	KY CALSNDSS	kt MKSGKEVE	LK NEGLEDIE	LM LNVCYDIINK
1	90 2	00 2	10 2	220 2	30 240
RMPFSPYI	CA GIGTDLIF	ME DAINHKAA	YQ GKLGFNYI	PIS PEANISMG	VH FHKVTNNEFR
	50 2	.60 . 2	70	280 2	90 300

Fig. 24B

VPVLLTAGGL APDNLFAIVK LSICHFGLEF GYRVSF....

10	20	30	40	50	60
ATGAATAATA			AACACAGTAT	TAGTATGCTT	ATTGTCATTA
70	80	. 90	100	110	120
CCTAATATAT		GGCCATAAAC	AATAACGCTA	AAAAGTACTA	CGGATTATAT
130	140	150	160	170	180
ATCAGTGGAC	AATATAAACC	CAGTGTTTCT	GTTTTCAGTA	ATTTTTCAGT	TAAAGAAACC
190	200	210	220	230	240
AATGTCATAA	CTAAAAACCT	TATAGCTTTA	AAAAAAGATG	TTGACTCTAT	TGAAACCAAG
250	260	270	280	290	300
ACTGATGCCA	GTGTAGGTAT	TAGTAACCCA	TCAAATTTTA	CTATCCCCTA	TACAGCTGTA
310	320	330	. 340	350	360
TTTCAAGATA	ATTCTGTCAA	TTTCAATGGA	ACTATTGGTT	ACACCTTTGC	TGAAGGTACA
370	380	390	400	410	420
AGAGTTGAAA	TAGAAGGTTC	TTATGAGGAA	TTTGATGTTA	AAAACCCTGG	AGGCTATACA
430	440	450	. 460	47.0	480
CTAAGTGATG	CCTATCGCTA	TTTTGCATTA	GCACGTGAAA	TGAAAGGTAA	TAGTTTTACA
490	500	510	520	530	540
CCTAAAGAAA	AAGTTTCTAA	TAGTATTTT	CACACTGTAA	TGAGAAATGA	TGGATTATCT
550	560	570			
ATAATATCTG	TTATAGTAAA	TGTTTGCTAC	GATTTCTCTT	TGAACAATTI	GTCAATATCG
610					
CCTTACATAT	GTGGAGGAGC	AGGGGTAGAT	GCTATAGAAT	TCTTCGATGI	ATTACACATT
670					
AAGTTTGCAT	ATCAAAGCAA	GCTAGGTATI	GCTTATTCT		CATTAGTCTC
730					
TTTGCTAGTI	TATATTACCA	TAAAGTAATO	GGCAATCAA!		AAATGTCCAA
790			-		
CATGTTGCT	AACTTGCAAG	TATACCTÁA			ACTTAATATT
850			-		_
GGTTATTTT	GAGGTGAAAT	TGGTGCAAG	A TTGACATTT	AA	

Fig. 25A

. 10	20	30	40	50	60
MNNKLKETII	NTVLVCLLSL	PNISSSKAIN	NNAKKYYGLY	ISGQYKPSVS	VESNESVKET
70	. 80	90	100	110	120
NVITKNLIAL	KKDVDSIETK	TDASVGISNP	SNFTIPYTAV	FQDNSVNFNG	TIGYTFAEGT
130	140	. 150	160	170	180
RVEIEGSYEE	FDVKNPGGYT	LSDAYRYFAL	AREMKGNSFT	PREKVSNSIF	HTVMRNDGLS
190	. 200	210	220	230	240
IISVIVNVCY	DESLNNLSIS	PYICGGAGVD	AIEFFDVLHI	KFAYQSKLGI	AYSLPSNISL
250	260	270	280	- 290	300
FASLYYHKVM	GNOFKNLNVO	HVAELASIPK	ITSAVATLNI	GYFGGEIGAR	LTF

Fig. 25B

10	20	30	40	50	60
ATGGCAAATT					
70	80	90	100	110	120
			AAAAATACAA		
130	140	150	160	170	180
TACATCAGTG	GACAGTATAA		TCTGTTTTTA		
190	200	210	220	230	240
ACCAATGTTC	ATACAGTACA	ACTCATGGCG	CTTAAAAAAG		
250	260	270	280	290	300
GATACTGGAA	ATAGCGCAGG	TATTAGCAAA	CCACAAAATT		
310	320	330	340	350	360
AAATTTCAAG	ATAATGTTGC	TGGTCTTAGC	GGTGCACTTG		
370	380	390	400	410	420
TTAAGGATTG	AAATGGGGTT	TTCTTATGAA	AAATTTGATG	CTAAAGACCT	TGGTGAGTAC
430	440	450	460	470	480
ACCAAAATAA	AAGATGCTTA	TAGATATTTT	GCTCTAGTAC	GTGAAATGCA	TGTTAGTCTC
490	500	510	520	530	540
ATTTATCCAA	AAGATAATAA	CACAGGAACA	CATTATACTG	TTATGAGAAA	TGATGGTATA
550	560	570	580	590	600
TCTATTTCTT	CTGCTACAGT	AAATGGCTGC	TATGATTCTT	TTTTCCAGTT	TATCTTTGTC
610	620	630	640	650	660
ACCTATATGT	GTATAGGCAT	CGGTATAGAT	GCTATAGAAT	TTCTTAATGC	ATACATATTA
670	680	690	700	710	720
AGTTTGCTTG	CCAAGGTAGT	TAAGGTGTTA	ACTTATTCTG	TATCTCCCAA	TGTTAATTTA
730	740	750	760	770	780
	GATATTATCA	TAAAGTGATG	GGCAATAAAT	TTAAAAATTT	ACCTGTTCAA
790	800	810	820		840
		GTATCCAAGA	GTTACATCTG	CAATTGCTAC	ACTTGATATT
850	860	870	880	890	
GGCTACCTCG	GTGGTGAAAT	TGGCATAAGA	TTTATATTTT	AA	• • • • • • • • • •
	•	Fig	. 26A		
		1.6	. 2011		
10	20	30	. 40	50	60
MYKKYKLMTA					NFSAKETNVH
70	80	•	100		
TVQLMALKKD					FFYSKGLRIE
130	140	150		170	
MGFSYEKFDA					
190	200	210		230	
ATVNGCYDSF	FQFIFVTYMC				SPNVNLFADG
	260	270			
YYHKVMGNKF	KNLPVQYVNT	LEEYPRVTSA	IATLDIGYLG		

Fig. 26B

10	20	30	40	50	60
ATGGGAAATT	CTATGAATAA	TAAAAGTCAA	TTCTTAATAA	GATTTATATT	TTTAACATGC
70	80	90	· 100	110	. 120
ATGCTGTCAT	TACCTAATAT	ATCTCTTTCA	AAAGTAAATA	ACGAAAAACA	TTCTGGTTTG
130	140	150	160	170	. 180
TATATTAGCG	GGCAATACAA	ACCCAGTGTT	TCTGTTTTCA	GTAATTTTTC	AGTTAAAGAA .
190	200	210	220	230	240
ACCAACTTTC	ATACAAAACA	TCTCATAGCT	CTTAAACAAG	ATGTTGATTC	TGTTGAAATT
250	260	270	280	290	300
GATACTGGTA	GTAATACAGC	AGGTATTAGT	AACCCATCTA	ACTTTACAAT	CCCTTATACT
310	320	330	340	350	360
GCAGAATTTC	AAGACAACCA	TACTAACTGC	AATGGCTCTA	TTGGTTATGC	TTTTGCTGAA
370	380	390	400	410	420
GGTCCAAGAA	TTGAAATAGA	ATTATCATAT	GAAAAATTTG	ATGTTAAAAA	TCCCACAGGG
430	440	450	460	. 47.0	480
TATACTACAG	TAAAAGATGC	TTATAGATAC	TTTGCTTTAG	CACGTGAAAT	AAATATTTCT
490	500	510	520	530	5 4.0
CTATTCCAAC	CAAAACAAAA	AGAAGGTAGT	GGAATTTACC	ATGTCGTAAT	GAAAAACGAT
550	560	570	580	590	600
GGGTTATCTA	TCTTATCCAA	TATAGTTAAT	ATTTGCTACG	ATTTTTCTTT	AAATAATTTA
610				•	
CCTATATCAC	CTTATTTATO	CGGAGGAAT	GGTATAAATO	CCATAGAATI	CTTTGACGCT
670					
TTACATGTG	A AATTTGCTT	A TCAAAGCAAG	G GCAGGAATT		ATTACGTAAA
730			•	-	
ATCAACTTA!	r TTATTGATG	r atattacta	C GAAGTAATA		TAAAAACCTG
790			-	_	
AAAGTCCAA	C ATGTACATG	A ACTTAAAGA	T AATCCAAAA		AGTTGCTACA
85		~	-	-	_
CTTGATATA	G CATATTTTG	G TAGTGAAGC	T GGCATAAGA	A TTATATTTT	A A

Fig. 27A

	∠∪	٥٥	40	50	60
MNNKSQFLIR	FIFLTCMLSL	PNISLSKVNN	EKHSGLYISG	QYKPSVSVFS	NFSVKETNFH
70	80	90	100	110	120
TKHLIALKQD	VDSVEIDTGS	NTAGISNPSN	FTIPYTAEFQ	DNHTNCNGSI	GYAFAEGPRI
130	140	150	160	.170	180
EIELSYEKFD	VKNPTGYTTV	KDAYRYFALA	REINISLFQP	KQKEGSGIYH	VVMKNDGLSI
190	200	210	220	230	240
LSNIVNICYD	FSLNNLPISP	YLCGGMGINA	IEFFDALHVK	FAYQSKAGIS	YQLLRKINLF
250	260	270	280	290	300
IDVYYYEVIS	NKFKNLKVQH	VHELKDNPKV	TSAVATLDIA	YFGSEAGIRI	IF

Fig. 27B

10	20	30	40	50	60
	— -		TGTACATCGT		ATTATCATCA
70	80	` 90	100	110	. 120
			AATAGTACAA	AACATTCTGG	ATTATATGTT
130	140	150	160	170	180
AGCGGACAAT	ATAAGCCCAG	CGTTTCCATT	TTTAGCAAAT	TTTCAGTAAA	AGAAACAAAT
190	200	210	220	230	240
ACACATACAG	TACAGTTAGT	AGCTCTTAAA	AAAGATGTTA	ATTCTATTTC	TATGAACATC
250	260	270	280	290	300
AGTAATGGTG	CTACAGGCAT	TAGCAAAGCA	ACAAATTTTA	ATCTTCCTTA	TGTTGCAGAA
310	320	. 330	. 340	. 350	360
TTTCAAGACA	ATGCCTTCAA	CTTCAGTGGA	GCTATTGGTT	ATTCACTTTT	
370	380	390	400	410	420
AACATTGAAG	TTGAAGGTTC	TTATGAAGAA	TTCGATGCCA	AAAATCCTGG	_
430	440	450	460	470	480
TTAAATGATG	CATTCCGCTA		GCACGTGAAA		. <u>.</u>
490	500	510	520	530	540
AATAAGCATC			GATATAAGTA		
550	560	570	-580	590	600
AGAAATAATG			ATGATAAATG		
610	620	630		650	. 660
			ACAGGAATAG		
670				•	TTACCAATTA
730					TGATCAATTT
TCAGACAACA					
					TACATCTGCA
850					
					CACACTTTAA
91(
		Fig	. 28A		
10	20	30	40	50	60
					FSKFSVKETN
		•		110	
					AIGYSLFEOL
130					
					DISKTYYTVM
. 190					
RNNGLSILSI	MINGCYNLPI	NDLSISPYFO	TGIGVDALE	FDALHLKLAI	QSKIGATYQL
250					
SDNISLFTNG	YYHQVIGDQE	KNLKVQYIGI	E LKENPKITSI	A VATLNVGYFO	GEIGVRLTL.

Fig. 28B

10	20	30	40	50	60
AAGCTTCTTA	TGAAGAATTT	GACGTTAAAA	ATCCTGAAGG	ATCTACTACA	GACTCCTATA
70	68	90	100	110	. 120
GATATTTCGC	GTTAGCACGT	GGCATGGATG	GTAATAATAT	TCCTACAAGT	CAAAAATTTA
130	140	150	160	170	180
CTGTAATGAG	AAACGACGGG	TTATTAATCT	CATCTGTTAT	GATAAATGGC	TGTTACAATG
190	200	210	220	230	240
TCATACTAAA	TGATATACAA	GCAGAACCTT	ACATATGTGC	AGGACTAGGA	GGAGATTTTA
250	260	270	280	290	300
TAGAATTCTT	CAATGGCTTT	CATGTTAAGC	TAGCTTATCA	AGGTAAAGTA	GGCATTAGTT
310	320	330	340	350	360
ATCAAATATT	CCCTGAAGTA	AGATTATTTA	TTGATGGATA	CTACCATAAA	GTAAAAGGCA
370	380	390	400	410	420
ACAAGTTTAA	AAATTTACAC	GTTCAACATG	TAGGTGCACT	TGCAGCACTC	CCTAAAGTTA
430	440	450	460	470	480
CATCTGCAGT	TGCAACACTT	AATATTGGAT	ACTTTGGTTG	TGAAGCTGGA	GTAAGATTCA
490	500	510	520	530	540
TATTTTAA					

Fig. 29A

60	5 0	40	, 30	20	10
SVMINGCYNV	VMRNDGLLIS	NNIPTSQKFT	YFALARGMDG	PEGSTTDSYR	ASYEEFDVKN
120	110	100	90	80	. 70
DGYYHKVKGN	QIFPEVRLFI	AYQGKVGISY	EFFNGFHVKL	ICAGLGGDFI	ILNDIQAEPY
180	. 170	160	150	140	130
	F	FGCEAGVRFI	SAVATLNIGY	GALAALPKVT	KFKNLHVQHV

Fig. 29B

10	20	30	40	50	60
ATGAATTATA	AGAAAATTCT	AGTAAGAAGC	GCGTTAATCT	CATTAATGTC	AATCTTACCA
70	80		100	110	. 120
TATCAGTCTT	TTGCAGATCC	TGTAGGTTCA	AGAACTAATG	ATAACAAAGA	AGGCTTCTAC
130	.140		160		180
ATTAGTGCAA	AGTACAATCC	AAGTATATCA	CACTTTAGAA	AATTCTCTGC	TGAAGAAACT
190	200	210	220	230	240
CCTATTAATG	GAACAAATTC	TCTCACTAAA	AAAGTTTTCG	GACTAAAGAA	AGATGGTGAT
250	260	270	280		
ATAACAAAAA	AAGACGATTT	TACAAGAGTA	GCTCCAGGCA	TTGATTTTCA	AAATAACTTA
310	320	330		• -	
ATATCAGGAT	TTTCAGGAAG	TATTGGTTAC	TCTATGGACG	GACCAAGAAT	AGAACTTGAA
370	380	390	400	410	420
GCTGCATATC	ACAATTTAAT	CCAAAAACAC	GATAACAATG	ATACTGATAA	TGGTGAATAC
430			460		
TATAAACATT	·TTGCATATCT	CGTAAAGATG	CCATGGAAGA	TCAGCCATAT	GTTGTTCTTA
490			520		
AAAATGACGG	CATAC				• • • • • • • • • •

Fig. 30A

	20	30	40	50	60
10 MNYKKILVRS	2U	VOCET DRUCS	PTNONKEGFY	ISAKYNPSIS	HFRKFSAEET
		90	100	110	120
70	80	JU		TSGFSGSIGY	SMDGPRIELE
PINGTNSLTK		ITKKDDFTRV	160	170	180
130	140	· 150			
AAVHNT.TOKH	DNNDTDNGEY	YKHFAYLVKM	PAKISHWIFT	KMTAI	•••••

Fig. 30B

		HV1	
OMP-LF CMP-LE CMP-LD OMP-LC OMP-LB P28	NONCREFFITT TLUSINSTEP GISTSDAUGH DRIVO-GN	.MVSASS HADADBKG GDI AQSANRTD NANSNDV.T.S. WKTPSC NTNEI.TEXD	90 89 90 89 94 64 91
MAP-1			•
OMP-1A OMP-1F OMP-10 OMP-1C OMP-1C OMP-1C OMP-1C OMP-1A	YSFKYDENIPF LGFAGAVGYL MNGPRIELEM SYETTEVRNO CHRYKNDAM: — KYVALTH: — RSGGKLSHAD DEFVFLINEY. . I.S. G. V.F.V. — — R.C. OQ — QURSCIPET S.Y.L. S. . I.S. G. F.V. — G. — R.C. DR — RASSYMATA SHYL. PALEFQ. LI S. S. SI. A. D. — A AYOK A. P. D. DT. SCDY Y. FG. SR — BADH. B. S.Y.V. . I.S. D. V.V. — R.C. SH — AADH. S. S R9. . I.S. D. V.V. — R.P. G. — H.C. — L.C. DTASSSTMAN TIS.NV. S. F.V. — R.P. G. — H.C. — L. DTASSSTMAN TIS.NV. HV3	: LLDISIMINA CYDVISPOIP	186 184 188 184 188 160 185
OMP-1F OMP-1S OMP-1O OMP-1C OMP-1B P28 HAP-1	I. IV	-A T	280 278 2 286 280 283 256 284 8L

Fig. 31

International application No. PCT/US98/19600

A. CLASSIFICATION OF SUBJECT MATTER IPC(6) : A01N 43/04; A61K 39/02 US CL : 514/44; 424/234.1					
According to International Patent Classification (IPC) or to both national classification and IPC					
	DS SEARCHED		•		
Minimum do	ocumentation searched (classification system followed	by classification symbols)			
U.S. : 514/44; 424/234.1					
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched					
Electronic d	ata base consulted during the international search (nar	ne of data base and, where practicable	, search terms used)		
APS, DIA	ALOG ms: erlichi?, protein?, antigen?, polypeptide?, dna, re	combinant?, clone?, dna, polynucleoti	de, nucleotide?		
C. DOC	UMENTS CONSIDERED TO BE RELEVANT				
Category*	Citation of document, with indication, where app	propriate, of the relevant passages	Relevant to claim No.		
A	US 5,789,176 A (DAWSON et al) 04 claims and entire document.	August 1998, see abstract,	1, 9, 11, 19, 21- 22		
A	US 5,401,656 A (DAWSON et al) 28 claims and entire document.	March 1995, see abstract,	1, 9, 11, 19, 21- 22		
A	US 5,413,931 A (DAWSON et al) (claims and entire document.	99 May 1995, see abstract,	1, 9, 11, 19, 21- 22		
Y,E	US 5,869,335 A (MUNDERLOH et abstract, claims and entire document.	al) 09 February 1999, see	1, 9		
X Furt	her documents are listed in the continuation of Box C	See patent family annex.	<u> </u>		
1	oocial categories of cited documents:	"T" later document published after the in date and not in conflict with the ap-			
	becoment defining the general state of the ert which is not considered be of particular relevance	the principle or theory underlying the	ne invention		
	rlier document published on or after the international filing date	"X" document of particular relevance; to considered novel or cannot be considered.			
ci	seument which may throw doubts on priority claim(s) or which is ted to establish the publication date of snother citation or other model consequences are sided).	when the document is taken eleme "Y" document of particular relevance; t	the claimed invention counce be		
•0• de	secial resson (as specified) comment referring to an oral disclosure, use, exhibition or other secs	considered to involve an invention combined with one or more other su being obvious to a person skilled in	re stop whon the document is such documents, such combination		
	ocument published prior to the intermstional filing date but later than a priority date claimed	*A* document member of the same pate			
Date of the	actual completion of the international search	Date of mailing of the international s	earch report		
18 FEBR	UARY 1999	25 FEB 1999			
Commissi	mailing address of the ISA/US oner of Patents and Trademarks	Authorized officer	WBC		
Box PCT Washingto	on, D.C. 20231	GINNY PORTNER	1 Var		
Facsimile l	No. (703) 305-3230	Telephone No. (703) 308-0196	V VV		

International application No. PCT/US98/19600

Box 1 Observations where certain claims were found unsearchable (Continuation of Item 1 of first sheet)
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. X Claims Nos.: 2-8, 10, 12-18, 20 because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically: The claims as submitted evidenced blank lines, therefore the claims were incomplete and found to be unscarchable.
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
·
1. As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on Protest The additional search fees were accompanied by the applicant's protest.
No protest accompanied the payment of additional search fees.

International application No.
PCT/US98/19600

	ution). DOCUMENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X 	Database Medline on Dialog, US National Library of Medicine (Bethesda, MD, USA,). BROUQUI, P. et al. 'Serologic diagnosis of	11,19, 21, 22
Y	human monocytic ehrlichiosis by immunoblot analysis'. Clinical Diagnostic Laboratory Immunology, November 1994, Vol. 1, No. 6, pages 645-649, see entire abstract.	1, 9
Y	Database Medline on Dialog, US National Library of Medicine (Bethesda, MD, USA,). BROUQUI, P. et al. 'Antigenic characterization of ehrlichiae: protein immunoblotting of Ehrlichia canis, Ehrlichia sennetsu, and Ehrlichia risticii'. Journal of Clinical Microbiology. May 1992, Vol. 30, No. 5, pages 1062-1066, see entire abstract.	19, 21, 22
X	Database Medline on Dialog, US National Library of Medicine	11, 21
	(Bethesda, MD, USA,). CHEN, SM et al. 'Identification of the	1
Y	antigenic constituents of Ehrlichia chaffeensis'. American Journal of Tropical Medicine and Hygiene. January 1994, Vol. 50, No. 1, page 52-58, see entire abstract.	
X .	Database Medline on Dialog, US National Library of Medicine	11, 19
- Y	(Bethesda, MD, USA,). CHEN, SHENG-MIN et al. 'Analysis and Ultrastructural localization of Ehrlichia chaffeensis proteins with monoclonal antibodies'. The American Journal of Tropical Medicine and hygiene. April 1996, Vol. 54, No. 4, pages 405-412, see entire abstract.	21, 22
Y,P	Database Medline on Dialog, US National Library of Medicine (Bethesda, MD, USA,). CHEN, SM et al. 'Western immunoblotting analysis of the antibody responses of patients with human monocytotropic ehrlichiosis to different strains of Ehrlichia chaffeensis and Ehrlichia canis'. Clinical Diagnostic and Laboratory Immunology. November 1997, Vol. 4, No. 6, pages 731-735, see entire abstract.	11, 19, 21, 22
Y	Database Medline on Dialog, US National Library of Medicine (Bethesda, MD, USA,). DAWSON, JE et al. 'The Interface between research and the diagnoses of an emerging tick-borne disease, human ehrlichiosis due to Ehrilichia chaffeensis'. Archives of Internal Medicine, 22 January 1996, Vol. 156, No. 2, pages 137-end, see entire document.	1
Y	Database Medline on Dialog, US National Library of Medicine (Bethesda, MD, USA,). KELLY, PJ et al. 'Serological evidence for antigenic relationships between Ehrlichia canis and Cowdria ruminantiu'. Research in Veterinary Science. March 1994, Vol. 56, No. 2, page 170-174, see entire abstract.	19

International application No. PCT/US98/19600

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
ζ - -	Database Medline on Dialog, US National Library of Medicine (Bethesda, MD, USA,). RIKIHISA, Y. et al. 'Enzyme linked immunosorbent assay and western immunoblot analyses of	19, 21, 22 9
	Ehrlichia- canis and canine granulocytic Ehrlichia infection'. Journal of Clinical Microbiology. January 1992, Vol. 30, No. 1, pages 143-148, see entire abstract.	
ď	Database Medline on Dialog, US National Library of Medicine (Bethesda, MD, USA,). YU, XJ et al. 'Sequence and characterization of an Ehrlichia chaffeensis gene encoding 314 amino acids highly homologous to the NAD A enzyme'. FEMS Microbiology Letters, 01 September 1997, Vol. 154, No. 1, pages 53-58, see entire document.	1, 9